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JC931 U.S. PTO
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10/25/00

JC931 U.S. PTO

Date: October 25, 2000
Docket No.: 2786-0140P

Assistant Commissioner for Patents
Box PATENT APPLICATION
Washington, D.C. 20231

Sir:

As authorized by the inventor(s), transmitted herewith for filing
is a patent application applied for on behalf of the inventor(s)
according to the provisions of 37 CFR 1.41(c).

Inventor(s): BERNSTEIN, Jeanne
LEVINE, Zurit

For: VARIANTS OF ALTERNATIVE SPLICING

Enclosed are:

- X A specification consisting of 61 pages
- X 48 sheet(s) of Formal drawings
- Certified copy of Priority Document(s)
- X Executed Declaration in accordance with 37 CFR 1.64 will follow
- A verified statement to establish small entity status under 37
CFR 1.9 and 37 CFR 1.27
- X Preliminary Amendment
- X Information Sheet
- Information Disclosure Statement, PTO-1449 with reference(s)

X Other Sequence Listing (52 pages)

The filing fee has been calculated as shown below:

LARGE ENTITY				SMALL ENTITY	
FOR	NO. FILED	NO. EXTRA	RATE FEE		RATE FEE
BASIC FEE	***** ***** *****	***** ***** *****	***** ***** \$710.00 *****	or	**** **** \$355.00 ****
TOTAL CLAIMS	43 - 20 =	23	x18 =\$ 414.00	or	x 9 = \$ 0.00
INDEPENDENT	1 - 3 =	0	x80 =\$ 0.00	or	x 40 = \$ 0.00
MULTIPLE DEPENDENT CLAIM PRESENTED <u>yes</u>			+270 = \$270.00	or	+135 = \$ 0.00
TOTAL \$1,394.00				TOTAL \$ 0.00	

X The application transmitted herewith is filed in accordance with 37 CFR 1.41(c). The undersigned has been authorized by the inventor(s) to file the present application. The original duly executed patent application together with the surcharge will be forwarded in due course.

X A check in the amount of \$1,394.00 to cover the filing fee and recording fee (if applicable) is enclosed.

____ The Government Filing Fee will be paid at the time of completion of the filing requirement.

____ Please charge Deposit Account No. 02-2448 in the amount of \$_____. A triplicate copy of this transmittal form is enclosed.

X Send Correspondence to: BIRCH, STEWART, KOLASCH & BIRCH, LLP
P. O. Box 747
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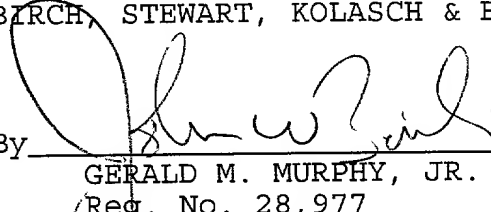
— No fee is enclosed.

If necessary, the Commissioner is hereby authorized in this, concurrent, and future replies, to charge payment or credit any overpayment to Deposit Account No. 02-2448 for any additional fees required under 37 C.F.R. 1.16 or under 37 C.F.R. 1.17; particularly, extension of time fees.

Respectfully submitted,

BIRCH, STEWART, KOLASCH & BIRCH, LLP

By

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IN THE U.S. PATENT AND TRADEMARK OFFICE

Applicant: BERNSTEIN, Jeanne et al.
Appl. No.: New Group:
Filed: October 25, 2000 Examiner:
For: VARIANTS OF ALTERNATIVE SPLICING

PRELIMINARY AMENDMENT

Assistant Commissioner for Patents
Washington, DC 20231

October 25, 2000

Sir:

The following preliminary amendments and remarks are respectfully submitted in connection with the above-identified application.

AMENDMENTS

IN THE CLAIMS:

Please amend the claims as follows:

Claim 5: Line 1, change "Claim 5" to --Claim 4--

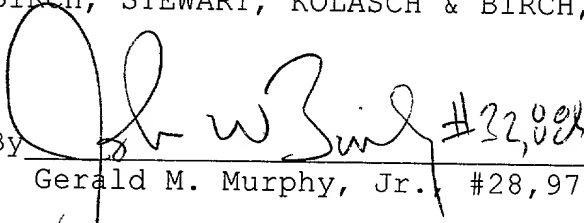
REMARKS

The amendment to the claims is merely to correct a typographical error and to place the application into better form for examination. Entry of the present amendment and favorable action on the above-identified application are respectfully requested.

If necessary, the Commissioner is hereby authorized in this, concurrent, and future replies, to charge payment or credit any overpayment to Deposit Account No. 02-2448 for any additional fees required under 37 C.F.R. § 1.16 or under 37 C.F.R. § 1.17; particularly, extension of time fees.

Respectfully submitted,

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VARIANTS OF ALTERNATIVE SPLICING

FIELD OF THE INVENTION

5 The present invention concerns novel nucleic acid sequences, vectors and host cells containing them, amino acid sequences encoded by said sequences, and antibodies reactive with said amino acid sequences, as well as pharmaceutical compositions comprising any of the above. The present invention further concerns methods for screening for candidate activators or deactivators utilizing said amino
10 acid sequences.

BACKGROUND OF THE INVENTION

Alternative splicing (AS) is an important regulatory mechanism in higher eukaryotes (P.A. Sharp, *Cell* 77, 805-8152 (1994). It is thought to be one of the most important mechanisms for differential expression related to tissue or
15 development stage specificity. It is known to play a major role in numerous biological systems, including human antibody responses, and sex determination in *Drosophila*, (S. Stamm, M.Q. Zhang, T.G. Marr and D.M. Helfman, *Nucleic Acids Research* 22, 1515-1526 (1994); B. Chabot, *Trends Genet.* 12, 472-478 (1996); R.E. Breitbart, A. Andreadis, B. Nadal-Ginard, *Annual Rev. Biochem.*,
20 56, 467-495 (1987); C.W. Smith, J.G. Patton, B. Nadal-Ginard, *Annu. Rev. Genet.*, 27, 527-577 (1989)).

Until recently it was commonly believed that alternative splicing existed in only a small fraction of genes (about 5%). A recent observation based on literature survey of known genes revises this conservative estimate to as high as
25 an estimate that at least 30% of human genes are alternatively spliced (M.S. Gelfand, I. Dubchak, I. Draluk and M. Zorn, *Nucleic Acids Research* 27, 301-302 (1999). The importance of the actual frequency of this phenomenon lies not only

in the direct impact on the number of proteins created (100,000 human genes, for example, would be translated to a much higher number of proteins), but also in the diversity of functionality derived from the process.

Several mechanisms at different stages may be held responsible for the complexity of higher eukaryote which include: alternative splicing at the transcription level, RNA editing at the post-transcriptional level, and post-translational modifications are the ones characterized to date.

GLOSSARY

10 In the following description and claims use will be made, at times, with a variety of terms, and the meaning of such terms as they should be construed in accordance with the invention is as follows:

"Variant nucleic acid sequence" – the sequence shown in any one of SEQ ID NO: 1 to SEQ ID NO: 26, sequences having at least 90% identity preferably 95% identity (see below) to said sequence and *fragments* (see below) of the above sequences of least 20 b.p. long. These sequences are sequences coding for a novel, naturally occurring, alternative splice variants of native and known genes. It should be emphasized that the novel variants of the present invention are naturally occurring sequences resulting from alternative splicing of genes and not merely truncated, mutated or fragmented forms of known sequences which are artificially produced.

"Variant product – also referred at times as the "variant protein" or "variant polypeptide" – is an amino acid sequence encoded by the variant nucleic acid sequence which is a naturally occurring mRNA sequence obtained as a result of alternative splicing. The amino acid sequence may be a peptide, a protein, as well as peptides or proteins having *chemically modified* amino acids (see below) such as a glycopeptide or glycoprotein. The variant products are shown in any one of SEQ ID NO: 27 to SEQ ID NO: 52. The term also includes *homology* (see below)

of said sequences in which one or more amino acids has been added, deleted, *substituted* (see below) or *chemically modified* (see below) as well as *fragments* (see below) of this sequence having at least 10 amino acids.

- 5 "Nucleic acid sequence" -- a sequence composed of DNA nucleotides, RNA nucleotides or a combination of both types and may includes natural nucleotides, chemically modified nucleotides and synthetic nucleotides.

- 10 "Amino acid sequence" -- a sequence composed of any one of the 20 naturally appearing amino acids, amino acids which have been *chemically modified* (see below), or composed of synthetic amino acids.

- 15 "Fragment of variant nucleic acid sequence" -- novel short stretch of nucleic acid sequences of at least 20 b.p., which does not appear as a continuous stretch in the *original nucleic acid sequence* (see below). The fragment may be a sequence which was previously undescribed in the context of the published RNA and which affects the amino acid sequence encoded by the known gene. For example, where the variant nucleic includes a sequence which was not included in the original sequence (for example a sequence which was an intron in the original sequence) the fragment may contain said additional sequence. The fragment may also be a region which is not an intron, which was not present in the original sequence. For example where the variant lacks a non-terminal region which was present in the original sequence. The two stretches of nucleotides spanning this region (upstream and downstream) are brought together by splicing
- 20 in the variant, but are spaced from each by the spliced out region in the original sequence and are thus not continuous in the original sequence. A continuous stretch of nucleic acids comprising said two splicing stretches of nucleotides is not present in the original sequence and thus falls under the definition of fragment.

"Fragments of variant products" - novel amino acid sequences coded by the *"fragment of variant nucleic acid sequence"* defined above.

5 *"Homologues of variants"* - amino acid sequences of variants in which one or more amino acids has been added, deleted or replaced. The addition, deletion or replacement should be in the regions or adjacent to regions where the variant differs from the *original sequence* (see below).

10 *"Conservative substitution"* - refers to the substitution of an amino acid in one class by an amino acid of the same class, where a class is defined by common physicochemical amino acid side chain properties and high substitution frequencies in homologous proteins found in nature, as determined, for example, by a standard Dayhoff frequency exchange matrix or BLOSUM matrix. [Six general classes of amino acid side chains have been categorized and include:
15 Class I (Cys); Class II (Ser, Thr, Pro, Ala, Gly); Class III (Asn, Asp, Gln, Glu); Class IV (His, Arg, Lys); Class V (Ile, Leu, Val, Met); and Class VI (Phe, Tyr, Trp). For example, substitution of an Asp for another class III residue such as Asn, Gln, or Glu, is a conservative substitution.

20 *"Non-conservative substitution"* - refers to the substitution of an amino acid in one class with an amino acid from another class; for example, substitution of an Ala, a class II residue, with a class III residue such as Asp, Asn, Glu, or Gln.

"Chemically modified" - when referring to the product of the invention, means a
25 product (protein) where at least one of its amino acid residues is modified either by natural processes, such as processing or other post-translational modifications, or by chemical modification techniques which are well known in the art. Among the numerous known modifications typical, but not exclusive examples include: acetylation, acylation, amidation, ADP-ribosylation, glycosylation, GPI anchor
30 formation, covalent attachment of a lipid or lipid derivative, methylation,

myristylation, pegylation, prenylation, phosphorylation, ubiquitination, or any similar process.

"Biologically active" - refers to the variant product having some sort of biological activity, for example, some physiologically measurable effect on target cells, molecules or tissues.

"Immunologically active" defines the capability of a natural, recombinant or synthetic variant product, or any fragment thereof, to induce a specific immune response in appropriate animals or cells and to bind with specific antibodies. Thus, for example, an immunologically active fragment of variant product denotes a fragment which retains some or all of the immunological properties of the variant product, e.g. can bind specific anti-variant product antibodies or which can elicit an immune response which will generate such antibodies or cause proliferation of specific immune cells which produce variant.

"Optimal alignment" - is defined as an alignment giving the highest percent identity score. Such alignment can be performed using a variety of commercially available sequence analysis programs, such as the local alignment program LALIGN using a ktup of 1, default parameters and the default PAM. A preferred alignment is the one performed using the CLUSTAL-W program from MacVector (TM), operated with an open gap penalty of 10.0, an extended gap penalty of 0.1, and a BLOSUM similarity matrix. If a gap needs to be inserted into a first sequence to optimally align it with a second sequence, the percent identity is calculated using only the residues that are paired with a corresponding amino acid residue (i.e., the calculation does not consider residues in the second sequences that are in the "gap" of the first sequence). In case of alignments of known gene sequences with that of the new variant, the optimal alignment invariably included aligning the identical parts of both sequences together, then

keeping apart and unaligned the sections of the sequences that differ one from the other.

"Having at least X% identity" - with respect to two amino acid or nucleic acid
5 sequence sequences, refers to the percentage of residues that are identical in the
two sequences when the sequences are optimally aligned. Thus, 90% amino acid
sequence identity means that 90% of the amino acids in two or more optimally
aligned polypeptide sequences are identical, however this definition explicitly
excludes sequences which are 100% identical with the original sequence from
10 which the variant of the invention was varied.

"Isolated nucleic acid molecule having an variant nucleic acid sequence" - is a
nucleic acid molecule that includes the coding variant nucleic acid sequence. Said
isolated nucleic acid molecule may include the variant nucleic acid sequence as
15 an independent insert; may include the variant nucleic acid sequence fused to an
additional coding sequences, encoding together a fusion protein in which the
variant coding sequence is the dominant coding sequence (for example, the
additional coding sequence may code for a signal peptide); the variant nucleic
acid sequence may be in combination with non-coding sequences, e.g., introns or
20 control elements, such as promoter and terminator elements or 5' and/or 3'
untranslated regions, effective for expression of the coding sequence in a suitable
host; or may be a vector in which the variant protein coding sequence is a
heterologous.

25 *"Expression vector"* - refers to vectors that have the ability to incorporate and
express heterologous DNA fragments in a foreign cell. Many prokaryotic and
eukaryotic expression vectors are known and/or commercially available.
Selection of appropriate expression vectors is within the knowledge of those
having skill in the art.

"Deletion" - is a change in either nucleotide or amino acid sequence in which one or more nucleotides or amino acid residues, respectively, are absent.

"Insertion" or "addition" - is that change in a nucleotide or amino acid
5 sequence which has resulted in the addition of one or more nucleotides or amino acid residues, respectively, as compared to the naturally occurring sequence.

"Substitution" - replacement of one or more nucleotides or amino acids by different nucleotides or amino acids, respectively. As regards amino acid
10 sequences the substitution may be conservative or non- conservative.

"Antibody" - refers to IgG, IgM, IgD, IgA, or IgG antibody. The definition includes polyclonal antibodies or monoclonal antibodies. This term refers to whole antibodies or fragments of the antibodies comprising the antigen-binding
15 domain of the anti-variant product antibodies, e.g. antibodies without the Fc portion, single chain antibodies, fragments consisting of essentially only the variable, antigen-binding domain of the antibody, etc.

"Distinguishing antibody" - an antibody capable of binding to the variant product
20 and not the original amino acid sequence from which it has been varied, or an antibody capable of binding to the original nucleic acid sequence and not to the variant production.

"Activator" - as used herein, refers to a molecule which mimics the effect of the
25 natural variant product or at times even increases or prolongs the duration of the biological activity of said product, as compared to that induced by the natural product. The mechanism may be by any mechanism known to prolonging activities of biological molecules such as binding to receptors; prolonging the lifetime of the molecules; increasing the activity of the molecules on its target;
30 increasing the affinity of molecules to its receptor; inhibiting degradation or

proteolysis of the molecules, or mimicking the biological activity of the variants on their targets, etc. Activators may be polypeptides, nucleic acids, carbohydrates, lipids, or derivatives thereof, or any other molecules which can bind to and activate the variant product.

5

"Deactivator" or (**"Inhibitor"**) - refers to a molecule which modulates the activity of the variant product in an opposite manner to that of the activator, by decreasing or shortening the duration of the biological activity of the variant product. This may be done by any mechanism known to deactivate or inhibit
10 biological molecules such as block of the receptor, block of active site, competition on binding site in target, enhancement of degradation, etc. Deactivators may be polypeptides, nucleic acids, carbohydrates, lipids, or derivatives thereof, or any other molecules which bind to and modulate the activity of said product.

15

"Treating a disease" - refers to administering a therapeutic substance effective to ameliorate symptoms associated with a disease, to lessen the severity or cure the disease, or to prevent the disease from occurring.

20 **"Detection"** – refers to a method of detection of a disease, disorder, pathological or normal condition. This term may refer to detection of a predisposition to a disease as well as for establishing the prognosis of the patient by determining the severity of the disease.

25 **"Probe"** – the variant nucleic acid sequence, or a sequence complementary therewith, when used to detect presence of other similar sequences in a sample. The detection is carried out by identification of hybridization complexes between the probe and the assayed sequence. The probe may be attached to a solid support or to a detectable label.

30

"Original sequence" – the amino acid or nucleic acid sequence from which the variant of the invention have been varied as a result of alternative slicing.

SUMMARY OF THE INVENTION

The present invention is based on the finding of several novel, naturally occurring splice variants, which are naturally occurring sequences obtained by alternative splicing of known genes. The novel splice variants of the invention are not merely truncated forms, fragments or mutations of known genes, but rather novel sequences which naturally occur within the body of individuals.

The term *"alternative splicing"* in the context of the present invention and claims refers to: intron inclusion, exon exclusion, addition or deletion of terminal sequences in the variant as compared to the original sequences, as well as to the possibility of *"intron retention"*. Intron retention is an intermediate stage in the processing of RNA transcripts, where prior to production of fully processed mRNA the intron (naturally spliced in the original sequence) is retained in the variant. These intermediately processed RNAs may have physiological significance and are also within the scope of the invention.

The novel variant products of the invention may have the same physiological activity as the original peptide from which they have been varied (although perhaps at a different level); may have an opposite physiological activity from the activity featured by the original peptide from which they are varied; may have a completely different, unrelated activity to the activity of the original from which they are varied; or alternatively may have no activity at all and this may lead to various diseases or pathological conditions. The variants may also differ from the original sequence by physiological properties not related to their activity such as their localization, their time and condition of expressing cleavage and degradation rates, association with other cellular or extracellular compounds and the like.

The novel variants may also serve for detection purposes, i.e. their presence or level may be indicative of a disease, disorder, pathological or normal condition or alternatively the ratio between the level variants and the level original peptide

from which they were varied, or the ratio to other variants may be indicative to a disease, disorder, pathological or normal condition.

For example, for detectional purposes, it is possible to establish differential expression of various variants in various tissues. A certain variant may be expressed mainly in one tissue, while the original sequence from which it has been varied, or another variant may, be expressed mainly in another tissue. Understanding of the distribution of the variants in various tissues may be helpful in basic research, for understanding the physiological function of the genes as well as may help in targeting pharmaceuticals or developing pharmaceuticals.

10 The study of the variants may also be helpful to distinguish various stages in the life cycles of the same type of cells which may also be helpful for development of pharmaceuticals for various pathological conditions in which cell cycles is non-normal, notably cancer.

Thus the detection may by determination of the presence or the level of expression of the variant within a specific cell population, comprising said presence or level between various cell types in a tissue, between different tissues and between individuals.

Thus the present invention provides by its first aspect, a novel isolated nucleic acid molecule comprising or consisting of any one of the coding sequence SEQ ID NO: 1 to SEQ ID NO: 26, fragments of said coding sequence having at least 20 nucleic acids (provided that said fragments are continuous stretches of nucleotides not present in the original sequence from which the variant was varied), or a molecule comprising a sequence having at least 90% identity, preferably 95% identity to SEQ ID NO: 1 to SEQ ID NO: 26, provided that the molecule is not completely identical to the original sequence from which the variant was varied.

The present invention further provides a protein or polypeptide comprising or consisting of an amino acid sequence encoded by any of the above nucleic acid sequences, termed herein "*variant product*", for example, an amino acid sequence having the sequence as depicted in any one of SEQ ID NO: 27 to SEQ ID NO: 52, fragments of the above amino acid sequence having a length of at least 10 amino

acids coded by the above fragments of the nucleic acid sequences, as well as homologues of the above amino acid sequences in which one or more of the amino acid residues has been substituted (by conservative or non-conservative substitution) added, deleted, or chemically modified.

5 The deletions, insertions and modifications should be in regions, or adjacent to regions, wherein the variant differs from the original sequence.

For example, where the variant is different from the original sequence by addition of a short stretch of 10 amino acids, in the terminal or non-terminal portion of the peptide, the invention also concerns homologues of that variant
10 where the additional short stretch is altered for example, it includes only 8 additional amino acids, includes 13 additional amino acids, or it includes 10 additional amino acids, however some of them being conservative or non-conservative substitutes of the original additional 10 amino acids of the novel variants. In all cases the changes in the homolog, as compared to the original
15 sequence, are in the same regions where the variant differs from the original sequence, or in regions adjacent to said region.

Another example is where the variant lacks a non-terminal region (for example of 20 amino acids) which is present in the original sequence (due for example to exon exclusion). The homologues may lack in the same region only 17
20 amino acids or 23 amino acids. Again the deletion is in the same region where the variant lacks a sequence as compared to the original sequence, or in a region adjacent thereto.

It should be appreciated that once a man versed in the art's attention is directed to the importance of a specific region, due to the fact that this region
25 differs in the variant as compared to the original sequence, there is no problem in derivating said specific region by addition to it, deleting from it, or substituting some amino acids in it. Thus homologues of variants which are derivated from the variant by changes (deletion, addition, substitution) only in said region as well as in regions adjacent to it are also a part of the present invention. Generally, if the
30 variant is distinguished from the original sequence by some sort of physiological

activity, then the homolog is distinguished from the original sequence in essentially the same manner.

The present invention further provides nucleic acid molecule comprising or consisting of a sequence which encodes the above amino acid sequences, 5 (including the fragments and homologues of the amino acid sequences). Due to the degenerative nature of the genetic code, a plurality of alternative nucleic acid sequences, beyond those depicted in any one of SEQ ID NO: 1 to SEQ ID NO: 26, can code for the amino acid sequence of the invention. Those alternative nucleic acid sequences which code for the same amino acid sequences codes by the 10 sequence SEQ ID NO: 1 to SEQ ID NO: 26 are also an aspect of the of the present invention.

The present invention further provides expression vectors and cloning vectors comprising any of the above nucleic acid sequences, as well as host cells transfected by said vectors.

15 The present invention still further provides pharmaceutical compositions comprising, as an active ingredient, said nucleic acid molecules, said expression vectors, or said protein or polypeptide.

These pharmaceutical compositions are suitable for the treatment of diseases and pathological conditions, which can be ameliorated or cured by raising the level 20 of any one of the variant products of the invention.

In the following a list will be given of diseases or pathological conditions which may be treated or detected by sequences of the invention (both nucleic acids, complementary and amino acids). As will be evident, the disease may be treated or detected also by antibodies which bind specifically to the amino acid sequences:

25 SEQ ID NO: 1 and sequences coded thereby, as well as antibodies, may be used for regulation (increase or decrease) of cytotoxicity induced by gram-negative bacteria as well as for detection of conditions of improper regulation. It may be used to combat existing infection by gram-negative bacteria.

SEQ ID NO: 2 and 18, sequences coded thereby and antibodies may be used for treatment and detection of diseases associated with faulty fat and vitamin absorption as well as for various digestive diseases.

SEQ ID NO: 3, sequences coded thereby and antibodies used for the
5 treatment and detection of disease involving faulty (too much or too little) blood coagulation, fibrinolysis or generation of bradykinn and angiotension. This may be used for treatment of disease of coagulation such as hemophilia or for treatment of excess coagulation or in arteriosclerosis.

SEQ ID NOS: 4, 5, 6, 7, 8, 23, 24, 25 and 26 sequences coded thereby and
10 antibodies cab be used for the treatment and detection of diseases or disorders associated with metalloproteinases (such as collogenases), and for the treatment and detection of diseases involved in erythrode progenitors.

SEQ ID NO: 9 and 19 sequences coded thereby and antibodies can be used for detection and treatment of thrombolytic disorders such as in pulmonary
15 embolism as well as diseases involved with the initiation of fibrinolysis.

SEQ ID NOS: 10 and 11 sequences coded thereby and antibodies can be used for the treatment and detection of diseases involved with fibrinolysis. The sequences and antibodies may act as regulators of fibronolysis.

SEQ ID NOS: 12, 20 and 21 and sequences coded thereby and antibodies
20 can be used for the detection and treatment of connective tissue irregularities and diseases involved with PDGF cell surface receptors.

SEQ ID NO: 13, 14 and 22, sequences encoded thereby and antibodies may be used for treatment of tumor, especially tumor due to mutation in tumor suppressor genes. In addition, may be used for detection of cancer or predisposition
25 to cancer. This gene may be associated to tumor in general and colorectal tumors in particular, as well as in to lymphatic and hematogenesis, metastasis of the esophageal squamous cell carcinoma, and may be used to detect or treat the disease.

SEQ ID NO: 15 sequences encoded thereby and antibodies, may be used for
30 the detection of treatment of diseases involved with the glutamate receptor in the

brain, such neurodegenerative disease, epilepsy, psychological disorders, brain tumors.

SEQ ID NOS: 16 and 17 sequences encoded thereby as well as antibodies, can be used for the treatment or detection of diseases involving regulation of
5 extracellular fluid volume mainly hypertension.

By a second aspect, the present invention provides a nucleic acid molecule comprising or consisting of a non-coding sequence which is complementary to that of any one of SEQ ID NO: 1 to SEQ ID NO: 26, or complementary to a sequence having at least 90% identity, preferably 95% identity to said sequence (with the
10 proviso added above) or a fragment of said two sequences (according to the above definition of fragment). The complementary sequence may be a DNA sequence which hybridizes with any one of SEQ of ID NO: 1 to SEQ ID NO: 26 or hybridizes to a portion of that sequence having a length sufficient to inhibit the transcription of the complementary sequence. The complementary sequence may be
15 a DNA sequence which can be transcribed into an mRNA being an antisense to the mRNA transcribed from any one of SEQ ID NO: 1 to SEQ ID NO: 26 or into an mRNA which is an antisense to a fragment of the mRNA transcribed from any one of SEQ ID NO: 1 to SEQ ID NO: 26 which has a length sufficient to hybridize with the mRNA transcribed from SEQ ID NO: 1 to SEQ ID NO: 26, so as to inhibit its
20 translation. The complementary sequence may also be the mRNA or the fragment of the mRNA itself.

The nucleic acids of the second aspect of the invention may be used for therapeutic or diagnostic applications for example as probes used for the detection of the variants of the invention. The presence of the variant transcript or the level of
25 the variant transcript may be indicative of a multitude of diseases, disorders and various pathological as well as normal conditions. In addition or alternatively, the ratio of the level of the transcripts of the variants of the invention may also be compared to that of the transcripts of the original sequences from which have been varied, or to the level of transcript of other variants, and said ratio may be

indicative to a multitude of diseases, disorders and various pathological and normal conditions.

The present invention also provides expression vectors comprising any one of the above defined complementary nucleic acid sequences and host cells
5 transfected with said nucleic acid sequences or vectors, being complementary to those specified in the first aspect of the invention.

The invention also provides anti-variant product antibodies, namely antibodies directed against the variant product which specifically bind to said variant product. Said antibodies are useful both for diagnostic and therapeutic
10 purposes. For example said antibody may be as an active ingredient in a pharmaceutical composition as will be explained below.

The present invention also provides pharmaceutical compositions comprising, as an active ingredient, the nucleic acid molecules which comprise or consist of said complementary sequences, or of a vector comprising said
15 complementary sequences. The pharmaceutical composition thus provides pharmaceutical compositions comprising, as an active ingredient, said anti-variant product antibodies.

The pharmaceutical compositions comprising said anti-variant product antibodies or the nucleic acid molecule comprising said complementary sequence,
20 are suitable for the treatment of diseases and pathological conditions where a therapeutically beneficial effect may be achieved by neutralizing the variant (either at the transcript or product level) or decreasing the amount of the variant product or blocking its binding to its target, for example, by the neutralizing effect of the antibodies, or by the effect of the antisense mRNA in decreasing the expression
25 level of the variant sequence.

According to the third aspect of the invention the present invention provides methods for detecting the level of the transcript (mRNA) of said variant product in a body fluid sample, or in a specific tissue sample, for example by use of probes comprising or consisting of said coding sequences; as well as methods for detecting
30 levels of expression of said product in tissue, e.g. by the use of antibodies capable

of specifically reacting with the variant products of the invention. Detection of the level of the expression of the variant of the invention in particular as compared to that of the original sequence from which it was varied or compared to other variant sequences all varied from the same original sequence may be indicative of a plurality of physiological or pathological conditions.

The method, according to this latter aspect, for detection of a nucleic acid sequence which encodes the variant product in a biological sample, comprises the steps of:

- (a) providing a probe comprising at least one of the nucleic acid sequences defined above;
- (b) contacting the biological sample with said probe under conditions allowing hybridization of nucleic acid sequences thereby enabling formation of hybridization complexes;
- (c) detecting hybridization complexes, wherein the presence of the complexes indicates the presence of nucleic acid sequence encoding the variant product in the biological sample.

The method as described above is qualitative, i.e. indicates whether the transcript is present in or absent from the sample. The method can also be quantitative, by determining the level of hybridization complexes and then calibrating said levels to determining levels of transcripts of the desired variant in the sample.

Both qualitative and quantitative determination methods can be used for diagnostic, prognostic and therapy planning purposes.

By a preferred embodiment the probe is part of a nucleic acid chip used for detection purposes, i.e. the probe is a part of an array of probes each present in a known location on a solid support.

The nucleic acid sequence used in the above method may be a DNA sequence an RNA sequence, etc; it may be a coding or a sequence or a sequence complementary thereto (for respective detection of RNA transcripts or coding-DNA sequences). By quantization of the level of hybridization complexes

and calibrating the quantified results it is possible also to detect the level of the transcript in the sample.

Methods for detecting mutations in the region coding for the variant product are also provided, which may be methods carried-out in a binary fashion, namely
5 merely detecting whether there is any mismatches between the normal variant nucleic acid sequence of the invention and the one present in the sample, or carried-out by specifically detecting the nature and location of the mutation.

The present invention also concerns a method for detecting variant product in a biological sample, comprising the steps of:

10 (a) contacting with said biological sample the antibody of the invention, thereby forming an antibody-antigen complex; and

(b) detecting said antibody-antigen complex

wherein the presence of said antibody-antigen complex correlates with the presence of variant product in said biological sample.

15 Many diseases are diagnosed by detecting the presence of antibodies against a protein characterizing the disease in the blood, serum or any other body fluid of the patient. The present invention also concerns a method for detecting anti-variant antibody in a biological sample, comprising:

(a) contacting said sample with the variant product of the invention,
20 thereby forming an antibody-antigen complex; and

(b) detecting said antibody-antigen complex

wherein the presence of said antibody-antigen complex correlates with the presence of anti-variant antibody in the sample.

As indicated above, both methods (for detection of variant product and for
25 detection of the anti-variant antibody) can be quantitized to determine the level or the amount of the variant or antibody in the sample, alone or in comparison to the level of the original amino acid sequence from which it was varied or compared to the level of antibodies against the original amino acid sequence, and qualitative and quantitative results may be used for diagnostic, prognostic and therapy planning
30 purposes.

The invention also concerns distinguishing antibodies, i.e. antibodies capable of binding either to the variant product or to the original sequence from which the variant has been varied, while not binding to the original sequence or the variant product respectively. These distinguishing antibodies may be used for
5 detection purposes.

By yet another aspect the invention also provides a method for identifying candidate compounds capable of binding to the variant product and modulating its activity (being either activators or deactivators). The method includes:

- (i) providing a protein or polypeptide comprising an amino acid
10 sequence substantially as depicted in any one of SEQ ID NO: 27 to 52, or a fragment of such a sequence;
- (ii) contacting a candidate compound with said amino acid sequence;
- (iii) measuring the physiological effect of said candidate compound on the activity of the amino acid sequences and selecting those compounds which
15 show a significant effect on said physiological activity.

The present invention also concerns compounds identified by the above methods described above, which compound may either be an activator of the variant product or a deactivator thereof.

20 BRIEF DESCRIPTION OF THE DRAWINGS

In order to understand the invention and to see how it may be carried out in practice, a preferred embodiment will now be described, by way of non-limiting example only, with reference to the accompanying drawings, in which:

Fig. 1 is a comparison between the amino acid sequence of SEQ ID NO: 27
25 and the original sequence from which it has been varied;

Fig. 2 is a comparison between the amino acid sequence of SEQ ID NO: 28 and the original sequence from which it has been varied;

Fig. 3 is a comparison between the amino acid sequence of SEQ ID NO: 29 and the original sequence from which it has been varied;

Fig. 4 is a comparison between the amino acid sequence of SEQ ID NO: 30 and the original sequence from which it has been varied;

Fig. 5 is a comparison between the amino acid sequence of SEQ ID NO: 31 and the original sequence from which it has been varied;

5 Fig. 6 is a comparison between the amino acid sequence of SEQ ID NO: 32 and the original sequence from which it has been varied;

Fig. 7 is a comparison between the amino acid sequence of SEQ ID NO: 33 and the original sequence from which it has been varied;

10 Fig. 8 is a comparison between the amino acid sequence of SEQ ID NO: 34 and the original sequence from which it has been varied;

Fig. 9 is a comparison between the amino acid sequence of SEQ ID NO: 35 and the original sequence from which it has been varied;

Fig. 10 is a comparison between the amino acid sequence of SEQ ID NO: 36 and the original sequence from which it has been varied;

15 Fig. 11 is a comparison between the amino acid sequence of SEQ ID NO: 37 and the original sequence from which it has been varied;

Fig. 12 is a comparison between the amino acid sequence of SEQ ID NO: 38 and the original sequence from which it has been varied;

20 Fig. 13 is a comparison between the amino acid sequence of SEQ ID NO: 39 and the original sequence from which it has been varied;

Fig. 14 is a comparison between the amino acid sequence of SEQ ID NO: 40 and the original sequence from which it has been varied;

Fig. 15 is a comparison between the amino acid sequence of SEQ ID NO: 41 and the original sequence from which it has been varied;

25 Fig. 16 is a comparison between the amino acid sequence of SEQ ID NO: 42 and the original sequence from which it has been varied;

Fig. 17 is a comparison between the amino acid sequence of SEQ ID NO: 43 and the original sequence from which it has been varied.

30 Fig. 18 is a comparison between the amino acid sequence of SEQ ID NO: 44 and the original sequence from which it has been varied.

Fig. 19 is a comparison between the amino acid sequence of SEQ ID NO: 45 and the original sequence from which it has been varied.

Fig. 20 is a comparison between the amino acid sequence of SEQ ID NO: 46 and the original sequence from which it has been varied.

5 Fig. 21 is a comparison between the amino acid sequence of SEQ ID NO: 47 and the original sequence from which it has been varied.

Fig. 22 is a comparison between the amino acid sequence of SEQ ID NO: 48 and the original sequence from which it has been varied.

10 Fig. 23 is a comparison between the amino acid sequence of SEQ ID NO: 49 and the original sequence from which it has been varied.

Fig. 24 is a comparison between the amino acid sequence of SEQ ID NO: 50 and the original sequence from which it has been varied.

Fig. 25 is a comparison between the amino acid sequence of SEQ ID NO: 51 and the original sequence from which it has been varied.

15 Fig. 26 is a comparison between the amino acid sequence of SEQ ID NO: 52 and the original sequence from which it has been varied.

DETAILED DESCRIPTION OF PREFERRED EMBODIMENTS

Example I: Comparison of variants with original sequences

20 Original sequences were obtained from GenBank Version 110. Comparison between the original sequences and the noval variant sequences was made using the Pileup application from the GCG suite version 10.0 (January 1999), with the default values:

Gap creation penalty (GapWeight): 8

25 Gap extension penalty (GapLengthWeight): 2

The comparison is shown in Fig. 1 to 26 which show the comparison of each of the variant products depicted in SEQ ID NO: 27 to 52 with the original sequence from which it was varied.

The following is a table that compares the sequences of the variants of the
30 invention to the original sequences from which they were varied and indicates here

the variant differs from the original sequence. The terminology NV-1 to NV-26 corresponds to SEQ ID NO: 1 to SEQ ID NO:26.

TABLE

5

Accession #	New Variant #	Description of the new variant
BPI_HUMAN	NV-1	Replacement of the last C-terminal 92 aa of the protein with alternative 100 aa, which has an additional potential transmembrane domain.
BAL_HUMAN	NV-2	Replacement of 121 C-terminal amino acids of the original protein by alternative 7 amino acids.
FA12_HUMAN	NV-3	Insertion of 30 amino acids in the catalytic domain of the original protein.
TIM1_HUMAN	NV-4	Replacement of 52 C-terminal amino acids of the original protein by alternative 9 amino acids. Missing 3 disulfide bonds.
TIM1_HUMAN	NV-5	Replacement of 94 C-terminal amino acids of the original protein by alternative 10 amino acids. Has only 1 out of 5 diS bonds, has an extra Cys in the added 10 amino acids.
TIM1_HUMAN	NV-6	Replacement of one of the exons of the original protein by a homologous exon.
TIM1_HUMAN	NV-7	Deletion of 47 amino acids of the original protein. Missing 2 disulfide bonds.
TIM1_HUMAN	NV-8	Deletion of 10 amino acids of the original protein. Missing one out of 2 glycosylation sites.
UROK_HUMAN	NV-9	Insertion of 62 amino acids in long A chain.
PAII_HUMAN	NV-10	Replacement of 140 C-terminal amino acids of the original protein, including the active site and 2 out of 3 glycosylation sites, by alternative 12 amino acids.
PAII_HUMAN	NV-11	Replacement of 17 C-terminal amino acids of the original protein by alternative 14 amino acids.
CTGF_HUMAN	NV-12	Deletion of 32 amino acids, at positions 152-185 of the original protein.
DCC_HUMAN	NV-13	Replacement of 1118 C-terminal amino acids of the original protein, including cytoplasmic domain, transmembrane region and part of the extracellular domain, by alternative 12 amino acids. The deleted region contains all 5 fibronectin type domains and 1 out of the 4 Ig-like C2 type domain.
DCC_HUMAN	NV-14	Replacement of 28 C-terminal amino acids from the cytoplasmic domain of the original protein by alternative 13 amino acids.
MGR4_HUMAN	NV-15	Deletion of 58 amino acids from the extracellular domain of the original protein.
MCR_HUMAN	NV-16	Replacement of 147 C-terminal amino acids, including part of the steroid binding domain of the original protein, by alternative 8 amino acids.

MCR_HUMAN	NV-17	Deletion of 147 C-terminal amino acids, including part of the steroid binding domain of the original protein.
BAL_HUMAN	NV-18	Deletion of last 102 aa at the C-terminus domain
UROK_HUMAN	NV-19	Replacement of 308 aa from C-terminus in 78 different aa
CTGF_HUMAN	NV-20	Deletion of N-terminus 276 aa
CTGF_HUMAN	NV-21	Replacement of C-terminus from the position 151 in 10 aa
DCC_HUMAN	NV-22	Deletion of C-terminus domain from aa 139
TIM1_HUMAN	NV-23	From amino-acid 103 of the original protein there is a replacement of the original C-terminus in 44 amino acids
TIM1_HUMAN	NV-24	Replacement of the first 67 amino acids of the original protein in different 13 amino acids, from amino acid 111 of the original protein there is a replacement of the original C-terminus in 25 amino acids
TIM1_HUMAN	NV-25	From amino acids 98 of the original protein there is a replacement of the original 109 amino acids of the C-terminus in 17 different amino-acids
TIM1_HUMAN	NV-26	Deletion of 65 amino acids of the N-terminus domain of the original protein.

Example II: Designation of the original sequences

Each novel variant of the invention is varied from an original sequence which has a known designation. The designation of the RNA sequences of the original sequences from which it was varied and the Accession Number of the original sequence are given below. First, information concerning the original sequence is given and then designation of the novel variants of the invention is given as NV-1 to NV-26 corresponding to SEQ ID NO: 1 to SEQ ID NO: 26. Diseases which can be treated or detected by the variants of the invention are also mentioned in the description of the original sequence.

BPI_HUMAN

BACTERICIDAL PERMEABILITY-INCREASING PROTEIN

FUNCTION: THE CYTOTOXIC ACTION OF BPI IS LIMITED TO MANY SPECIES OF GRAM-NEGATIVE BACTERIA; THIS SPECIFICITY MAY BE

EXPLAINED BY A STRONG AFFINITY OF THE VERY BASIC N-TERMINAL HALF FOR THE NEGATIVELY CHARGED LIPOPOLYSACCHARIDES THAT ARE UNIQUE TO THE GRAM-NEGATIVE BACTERIAL OUTER ENVELOPE.

5 SUBCELLULAR LOCATION: MEMBRANE-ASSOCIATED IN POLYMORPHONUCLEAR LEUKOCYTES (PMN) GRANULES.

TISSUE SPECIFICITY: RESTRICTED TO CELLS OF THE MYELOID SERIES.

10 DOMAIN: THE N-TERMINAL REGION MAY BE EXPOSED TO THE INTERIOR OF THE GRANULE, WHEREAS THE C-TERMINAL PORTION MAY BE EMBEDDED IN THE MEMBRANE. DURING PHAGOCYTOSIS AND DEGRANULATION, PROTEASES MAY BE RELEASED AND ACTIVATED AND CLEAVE BPI AT THE JUNCTION OF THE N- AND C-TERMINAL PORTIONS OF THE MOLECULE, PROVIDING
15 CONTROLLED RELEASE OF THE N-TERMINAL ANTIBACTERIAL FRAGMENT WHEN BACTERIA ARE INGESTED.

20 NV-1:

BAL_HUMAN

25 BILE-SALT-ACTIVATED LIPASE (BAL) CHOLESTEROL ESTERASE

FUNCTION: CATALYZES FAT AND VITAMIN ABSORPTION. ACTS IN CONCERT WITH PANCREATIC LIPASE AND COLIPASE FOR THE COMPLETE DIGESTION OF DIETARY TRIGLYCERIDES.

30 CATALYTIC ACTIVITY: TRIACYLGLYCEROL + H(2)O = DIACYLGLYCEROL + A FATTY ACID ANION.

CATALYTIC ACTIVITY: A STERYL ESTER + H(2)O = A STEROL + A FATTY ACID.

35 ENZYME REGULATION: ACTIVATED BY BILE SALTS CONTAINING A 7-HYDROXYL GROUP.

TISSUE SPECIFICITY: MAMMARY GLAND, AND PANCREAS.

SIMILARITY: BELONGS TO THE TYPE-B CARBOXYLESTERASE/LIPASE FAMILY.

40 NV-2: NV_18

FA12_HUMAN

COAGULATION FACTOR XII

5 FUNCTION: FACTOR XII IS A SERUM GLYCOPROTEIN THAT PARTICIPATES IN THE INITIATION OF BLOOD COAGULATION, FIBRINOLYSIS, AND THE GENERATION OF BRADYKININ AND ANGIOTENSIN.

10 CATALYTIC ACTIVITY: CLEAVES SELECTIVELY ARG-|-ILE BONDS AND ACTIVATES COAGULATION FACTORS VII AND XI.

PTM: O- AND N-GLYCOSYLATED.

DISEASE: DEFECTS IN F12 DO NOT CAUSE ANY CLINICAL SYMPTOMS. THE SOLE EFFECT IS THAT WHOLE-BLOOD CLOTTING TIME IS PROLONGED.

15 MISCELLANEOUS: FACTOR XII, PREKALLIKREIN, AND HMW KININOGEN FORM A COMPLEX BOUND TO AN ANIONIC SURFACE. PREKALLIKREIN IS CLEAVED BY FACTOR XII TO FORM KALLIKREIN, WHICH THEN CLEAVES FACTOR XII FIRST TO ALPHA-FACTOR XIIA AND THEN TO BETA-FACTOR XIIA. ALPHA-FACTOR XIIA ACTIVATES
20 FACTOR XI TO FACTOR XIA.

NV-3:

25

TIM1_HUMAN

METALLOPROTEINASE INHIBITOR 1

30 FUNCTION: COMPLEXES WITH METALLOPROTEINASES (SUCH AS COLLAGENASES) AND IRREVERSIBLY INACTIVATE THEM. ALSO MEDIATES ERYTHROPOIESIS IN VITRO; BUT, UNLIKE IL-3, IT IS SPECIES-SPECIFIC, STIMULATING THE GROWTH AND DIFFERENTIATION OF ONLY HUMAN AND MURINE ERYTHROID PROGENITORS.

35 PTM: THE ACTIVITY OF TIMP-1 IS DEPENDENT ON THE PRESENCE OF DISULFIDE BONDS.

SIMILARITY: BELONGS TO THE TIMP FAMILY.

40 NV-4; NV_23, NV_24, NV_24, NV_26

TIM1_HUMAN

METALLOPROTEINASE INHIBITOR 1

5

NV-5:

TIM1_HUMAN

10

METALLOPROTEINASE INHIBITOR 1

NV-6:

15

TIM1_HUMAN

METALLOPROTEINASE INHIBITOR 1

20

NV-7

TIM1_HUMAN

25

METALLOPROTEINASE INHIBITOR 1

NV-8

30

UROK_HUMAN

UROKINASE-TYPE PLASMINOGEN ACTIVATOR

35

FUNCTION: POTENT PLASMINOGEN ACTIVATOR AND IS CLINICALLY USED FOR THERAPY OF THROMBOLYTIC DISORDERS.

40 CATALYTIC ACTIVITY: SPECIFIC CLEAVAGE OF ARG-|-VAL BOND IN PLASMINOGEN TO FORM PLASMIN.

SUBUNIT: FOUND IN HIGH AND LOW MOLECULAR MASS FORMS. EACH CONSISTS OF TWO CHAINS, A AND B. THE HIGH MOLECULAR MASS FORM CONTAINS A LONG CHAIN A. CLEAVAGE OCCURS AFTER RESIDUE 155 IN THE LOW MOLECULAR MASS FORM TO YIELD
45 A SHORT A1 CHAIN.

PHARMACEUTICAL: AVAILABLE UNDER THE NAME ABBOKINASE (ABBOTT). USED IN PULMONARY EMBOLISM (PE) TO INITIATES FIBRINOLYSIS.

5 NV-9: NV_19

PAI1_HUMAN

10 PLASMINOGEN ACTIVATOR INHIBITOR-1, ENDOTHELIAL

FUNCTION: THIS INHIBITOR ACTS AS "BAIT" FOR TISSUE PLASMINOGEN ACTIVATOR, UROKINASE, AND PROTEIN C. ITS RAPID INTERACTION WITH TPA MAY FUNCTION AS A MAJOR CONTROL POINT IN THE REGULATION OF FIBRINOLYSIS.

15 DISEASE: HIGH CONCENTRATIONS OF THIS PROTEIN HAVE BEEN ASSOCIATED WITH HUMAN THROMBOEMBOLIC DISEASE.

MISCELLANEOUS: PAI1 IS INACTIVATED BY PROTEOLYTIC ATTACK OF THE UROKINASE-TYPE (U-PA) AND THE TISSUE-TYPE (TPA),
20 CLEAVING THE 369(R)-370(M) BOND.

NV-10:

PAI1_HUMAN

25

PLASMINOGEN ACTIVATOR INHIBITOR-1, ENDOTHELIAL

NV-11:

30

CTGF_HUMAN

CONNECTIVE TISSUE GROWTH FACTOR

35 FUNCTION: MAJOR CONNECTIVE TISSUE MITOATTRACTANT SECRETED BY HUMAN VASCULAR ENDOTHELIAL CELLS. THIS IMMEDIATE-EARLY PROTEIN MAY BIND ONE OF THE PDGF CELL SURFACE RECEPTORS.

SUBUNIT: MONOMER.

40 ALTERNATIVE PRODUCTS: A SHORTER FORM MAY BE PRODUCED BY ALTERNATIVE SPLICING OF THE SAME GENE: VARSPLIC 172 198 MISSING (IN SHORT FORM).

SIMILARITY: BELONGS TO THE INSULIN-LIKE GROWTH FACTOR BINDING PROTEIN FAMILY. CEF-10/CYR61/CTFG/FISP-12/NOV

NV-12; NV_20, NV_21

5

DCC_HUMAN

TUMOR SUPPRESSOR PROTEIN DCC

FUNCTION: IMPLICATED AS A TUMOR SUPPRESSOR GENE.

10 SUBCELLULAR LOCATION: TYPE I MEMBRANE PROTEIN.

TISSUE SPECIFICITY: FOUND IN AXONS OF THE CENTRAL AND PERIPHERAL NERVOUS SYSTEM AND IN DIFFERENTIATED CELL TYPES OF THE INTESTINE.

DISEASE: COLORECTAL TUMORS THAT LOST THEIR CAPACITY TO
15 DIFFERENTIATE INTO MUCUS PRODUCING CELLS UNIFORMLY LACK DCC EXPRESSION. INACTIVATION OF DCC DUE TO ALLELIC DELETION AND/OR POINT MUTATIONS MAY CAUSE BOTH LYMPHATIC AND HEMATOGENOUS METASTASIS OF OESOPHAGEAL SQUAMOUS CELL CARCINOMAS.

20 SIMILARITY: CONTAINS 4 IMMUNOGLOBULIN-LIKE C2-TYPE DOMAINS.

SIMILARITY: CONTAINS 6 FIBRONECTIN TYPE III-LIKE DOMAINS.

25 NV-13; NV_22

DCC_HUMAN

30

TUMOR SUPPRESSOR PROTEIN DCC

NV-14:

35

MGR4_HUMAN

METABOTROPIC GLUTAMATE RECEPTOR 4

5 **FUNCTION:** RECEPTOR FOR GLUTAMATE. THE ACTIVITY OF THIS RECEPTOR IS MEDIATED BY A G-PROTEIN THAT INHIBITS ADENYLATE CYCLASE ACTIVITY.

SUBCELLULAR LOCATION: INTEGRAL MEMBRANE PROTEIN.

10 **TISSUE SPECIFICITY:** STRONGLY EXPRESSED IN THE CEREBELLUM. EXPRESSED AT LOW LEVELS IN HIPPOCAMPUS, HYPOTHALAMUS AND THALAMUS. NO EXPRESSION DETECTED IN LIVER.

SIMILARITY: BELONGS TO FAMILY 3 OF G-PROTEIN COUPLED RECEPTORS. STRONGEST, TO MGLUR6.

15 **NV-15:**

MCR_HUMAN

20 **MINERALOCORTICOID RECEPTOR**

FUNCTION: RECEPTOR FOR BOTH MINERALOCORTICIDS (MC) SUCH AS ALDOSTERONE AND GLUCOCORTICIDS (GC) SUCH AS CORTICOSTERONE OR CORTISOL. THE EFFECT OF MC IS TO INCREASE ION AND WATER TRANSPORT AND THUS RAISE EXTRACELLULAR FLUID VOLUME AND BLOOD PRESSURE AND LOWER POTASSIUM LEVELS.

SUBCELLULAR LOCATION: NUCLEAR.

30 **DOMAIN:** COMPOSED OF THREE DOMAINS: A MODULATING N-TERMINAL DOMAIN, A DNA-BINDING DOMAIN AND A C-TERMINAL STEROID-BINDING DOMAIN.

SIMILARITY: BELONGS TO THE NUCLEAR HORMONE RECEPTORS FAMILY. NR3 SUBFAMILY.

35 **NV-16:**

MCR_HUMAN

40 **MINERALOCORTICOID RECEPTOR**

45 **NV-17:**

Example III: Variant nucleic acid sequence

The nucleic acid sequences of the invention include nucleic acid sequences which encode variant product and fragments and analogs thereof. The
5 nucleic acid sequences may alternatively be sequences complementary to the above coding sequence, or to a region of said coding sequence. The length of the complementary sequence is sufficient to avoid the expression of the coding sequence. The nucleic acid sequences may be in the form of RNA or in the form of DNA, and include messenger RNA, synthetic RNA and DNA, cDNA, and
10 genomic DNA. The DNA may be double-stranded or single-stranded, and if single-stranded may be the coding strand or the non-coding (anti-sense, complementary) strand. The nucleic acid sequences may also both include dNTPs, rNTPs as well as non naturally occurring sequences. The sequence may also be a part of a hybrid between an amino acid sequence and a nucleic acid
15 sequence.

In a general embodiment, the nucleic acid sequence has at least 90% identity, preferably 95% identity with any one of the sequence identified as SEQ ID NO: 1 to SEQ ID NO: 26 provided that this sequence is not completely identical with that of the original sequence.

20 The nucleic acid sequences may include the coding sequence by itself. By another alternative the coding region may be in combination with additional coding sequences, such as those coding for fusion protein or signal peptides, in combination with non-coding sequences, such as introns and control elements, promoter and terminator elements or 5' and/or 3' untranslated regions, effective
25 for expression of the coding sequence in a suitable host, and/or in a vector or host environment in which the variant nucleic acid sequence is introduced as a heterologous sequence.

The nucleic acid sequences of the present invention may also have the product coding sequence fused in-frame to a marker sequence which allows for
30 purification of the variant product. The marker sequence may be, for example, a

hexahistidine tag to provide for purification of the mature polypeptide fused to the marker in the case of a bacterial host, or, the marker sequence may be a hemagglutinin (HA) tag when a mammalian host, e.g. COS-7 cells, is used. The HA tag corresponds to an epitope derived from the influenza hemagglutinin protein (Wilson, I., *et al. Cell* 37:767 (1984)).

Also included in the scope of the invention are fragments as defined above also referred to herein as oligonucleotides, typically having at least 20 bases, preferably 20-30 bases corresponding to a region of the coding-sequence nucleic acid sequence. The fragments may be used as probes, primers, and when complementary also as antisense agents, and the like, according to known methods.

As indicated above, the nucleic acid sequence may be substantially a depicted in any one of SEQ ID NO: 1 to SEQ ID NO: 26 or fragments thereof or sequences having at least 90% identity, preferably 95% identity to the above sequence as explained above. Alternatively, due to the degenerative nature of the genetic code, the sequence may be a sequence coding for any one of the amino acid sequence of SEQ ID NO: 27 to SEQ ID NO: 52, or fragments or analogs of said amino acid sequence.

20 A. Preparation of nucleic acid sequences

The nucleic acid sequences may be obtained by screening cDNA libraries using oligonucleotide probes which can hybridize to or PCR-amplify nucleic acid sequences which encode the variant products disclosed above. cDNA libraries prepared from a variety of tissues are commercially available and procedures for screening and isolating cDNA clones are well-known to those of skill in the art. Such techniques are described in, for example, Sambrook *et al.* (1989) *Molecular Cloning: A Laboratory Manual* (2nd Edition), Cold Spring Harbor Press, Plainview, N.Y. and Ausubel *et al.* (1989) *Current Protocols in Molecular Biology*, John Wiley & Sons, New York, N.Y.

The nucleic acid sequences may be extended to obtain upstream and downstream sequences such as promoters, regulatory elements, and 5' and 3' untranslated regions (UTRs). Extension of the available transcript sequence may be performed by numerous methods known to those of skill in the art, such as
5 PCR or primer extension (Sambrook *et al.*, *supra*), or by the RACE method using, for example, the Marathon RACE kit (Clontech, Cat. # K1802-1).

Alternatively, the technique of "restriction-site" PCR (Gobinda *et al.* *PCR Methods Applic.* 2:318-22, (1993)), which uses universal primers to retrieve flanking sequence adjacent a known locus, may be employed. First, genomic
10 DNA is amplified in the presence of primer to a linker sequence and a primer specific to the known region. The amplified sequences are subjected to a second round of PCR with the same linker primer and another specific primer internal to the first one. Products of each round of PCR are transcribed with an appropriate RNA polymerase and sequenced using reverse transcriptase.

15 Inverse PCR can be used to amplify or extend sequences using divergent primers based on a known region (Triglia, T. *et al.*, *Nucleic Acids Res.* 16:8186, (1988)). The primers may be designed using OLIGO(R) 4.06 Primer Analysis Software (1992; National Biosciences Inc, Plymouth, Minn.), or another appropriate program, to be 22-30 nucleotides in length, to have a GC content of
20 50% or more, and to anneal to the target sequence at temperatures about 68-72°C.

The method uses several restriction enzymes to generate a suitable fragment in the known region of a gene. The fragment is then circularized by intramolecular ligation and used as a PCR template.

Capture PCR (Lagerstrom, M. *et al.*, *PCR Methods Applic.* 1:111-19, 25 (1991)) is a method for PCR amplification of DNA fragments adjacent to a known sequence in human and yeast artificial chromosome DNA. Capture PCR also requires multiple restriction enzyme digestions and ligations to place an engineered double-stranded sequence into a flanking part of the DNA molecule before PCR.

Another method which may be used to retrieve flanking sequences is that of Parker, J.D., *et al.*, *Nucleic Acids Res.*, **19**:3055-60, (1991)). Additionally, one can use PCR, nested primers and PromoterFinder™ libraries to "walk in" genomic DNA (PromoterFinder™; Clontech, Palo Alto, CA). This process avoids the need
5 to screen libraries and is useful in finding intron/exon junctions. Preferred libraries for screening for full length cDNAs are ones that have been size-selected to include larger cDNAs. Also, random primed libraries are preferred in that they will contain more sequences which contain the 5' and upstream regions of genes.

A randomly primed library may be particularly useful if an oligo d(T)
10 library does not yield a full-length cDNA. Genomic libraries are useful for extension into the 5' nontranslated regulatory region.

The nucleic acid sequences and oligonucleotides of the invention can also be prepared by solid-phase methods, according to known synthetic methods. Typically, fragments of up to about 100 bases are individually synthesized, then
15 joined to form continuous sequences up to several hundred bases.

B. Use of variant nucleic acid sequence for the production of variant products

20 In accordance with the present invention, nucleic acid sequences specified above may be used as recombinant DNA molecules that direct the expression of variant products.

As will be understood by those of skill in the art, it may be advantageous to produce variant product-encoding nucleotide sequences possessing codons
25 other than those which appear in any one of SEQ ID NO: 1 to SEQ ID NO: 26 which are those which naturally occur in the human genome. Codons preferred by a particular prokaryotic or eukaryotic host (Murray, E. *et al.* *Nuc Acids Res.*, **17**:477-508, (1989)) can be selected, for example, to increase the rate of variant product expression or to produce recombinant RNA transcripts having desirable

properties, such as a longer half-life, than transcripts produced from naturally occurring sequence.

The nucleic acid sequences of the present invention can be engineered in order to alter a variant product coding sequence for a variety of reasons, including but not limited to, alterations which modify the cloning, processing
5 and/or expression of the product. For example, alterations may be introduced using techniques which are well known in the art, e.g., site-directed mutagenesis, to insert new restriction sites, to alter glycosylation patterns, to change codon preference, etc.

10 The present invention also includes recombinant constructs comprising one or more of the sequences as broadly described above. The constructs comprise a vector, such as a plasmid or viral vector, into which a nucleic acid sequence of the invention has been inserted, in a forward or reverse orientation. In a preferred aspect of this embodiment, the construct further comprises
15 regulatory sequences, including, for example, a promoter, operably linked to the sequence. Large numbers of suitable vectors and promoters are known to those of skill in the art, and are commercially available. Appropriate cloning and expression vectors for use with prokaryotic and eukaryotic hosts are also described in Sambrook, *et al.*, (*supra*).

20 The present invention also relates to host cells which are genetically engineered with vectors of the invention, and the production of the product of the invention by recombinant techniques. Host cells are genetically engineered (i.e., transduced, transformed or transfected) with the vectors of this invention which may be, for example, a cloning vector or an expression vector. The vector may
25 be, for example, in the form of a plasmid, a viral particle, a phage, etc. The engineered host cells can be cultured in conventional nutrient media modified as appropriate for activating promoters, selecting transformants or amplifying the expression of the variant nucleic acid sequence. The culture conditions, such as temperature, pH and the like, are those previously used with the host cell selected
30 for expression, and will be apparent to those skilled in the art.

The nucleic acid sequences of the present invention may be included in any one of a variety of expression vectors for expressing a product. Such vectors include chromosomal, nonchromosomal and synthetic DNA sequences, e.g., derivatives of SV40; bacterial plasmids; phage DNA; baculovirus; yeast
5 plasmids; vectors derived from combinations of plasmids and phage DNA, viral DNA such as vaccinia, adenovirus, fowl pox virus, and pseudorabies. However, any other vector may be used as long as it is replicable and viable in the host. The appropriate DNA sequence may be inserted into the vector by a variety of procedures. In general, the DNA sequence is inserted into an appropriate
10 restriction endonuclease site(s) by procedures known in the art. Such procedures and related sub-cloning procedures are deemed to be within the scope of those skilled in the art.

The DNA sequence in the expression vector is operatively linked to an appropriate transcription control sequence (promoter) to direct mRNA synthesis.
15 Examples of such promoters include: LTR or SV40 promoter, the *E.coli lac* or *trp* promoter, the phage lambda *PL* promoter, and other promoters known to control expression of genes in prokaryotic or eukaryotic cells or their viruses. The expression vector also contains a ribosome binding site for translation initiation, and a transcription terminator. The vector may also include
20 appropriate sequences for amplifying expression. In addition, the expression vectors preferably contain one or more selectable marker genes to provide a phenotypic trait for selection of transformed host cells such as dihydrofolate reductase or neomycin resistance for eukaryotic cell culture, or such as tetracycline or ampicillin resistance in *E.coli*.

25 The vector containing the appropriate DNA sequence as described above, as well as an appropriate promoter or control sequence, may be employed to transform an appropriate host to permit the host to express the protein. Examples of appropriate expression hosts include: bacterial cells, such as *E.coli*, *Streptomyces*, *Salmonella typhimurium*; fungal cells, such as yeast; insect cells
30 such as *Drosophila* and *Spodoptera Sf9*; animal cells such as CHO, COS, HEK

293 or Bowes melanoma; adenoviruses; plant cells, etc. The selection of an appropriate host is deemed to be within the scope of those skilled in the art from the teachings herein. The invention is not limited by the host cells employed.

In bacterial systems, a number of expression vectors may be selected
5 depending upon the use intended for the variant product. For example, when large quantities of variant product are needed for the induction of antibodies, vectors which direct high level expression of fusion proteins that are readily purified may be desirable. Such vectors include, but are not limited to, multifunctional *E.coli* cloning and expression vectors such as *Bluescript*(R)
10 (Stratagene), in which the variant polypeptide coding sequence may be ligated into the vector in-frame with sequences for the amino-terminal Met and the subsequent 7 residues of beta-galactosidase so that a hybrid protein is produced; *pIN* vectors (Van Heeke & Schuster *J. Biol. Chem.* 264:5503-5509, (1989)); *pET* vectors (Novagen, Madison WI); and the like.

15 In the yeast *Saccharomyces cerevisiae* a number of vectors containing constitutive or inducible promoters such as alpha factor, alcohol oxidase and PGH may be used. For reviews, see Ausubel *et al.* (*supra*) and Grant *et al.*, (*Methods in Enzymology* 153:516-544, (1987)).

In cases where plant expression vectors are used, the expression of a
20 sequence encoding variant product may be driven by any of a number of promoters. For example, viral promoters such as the 35S and 19S promoters of *CaMV* (Brisson *et al.*, *Nature* 310:511-514, (1984)) may be used alone or in combination with the omega leader sequence from TMV (Takamatsu *et al.*, *EMBO J.*, 6:307-311, (1987)). Alternatively, plant promoters such as the small
25 subunit of RUBISCO (Coruzzi *et al.*, *EMBO J.* 3:1671-1680, (1984); Broglie *et al.*, *Science* 224:838-843, (1984)); or heat shock promoters (Winter J and Sinibaldi R.M., *Results Probl. Cell Differ.*, 17:85-105, (1991)) may be used. These constructs can be introduced into plant cells by direct DNA transformation or pathogen-mediated transfection. For reviews of such techniques, see Hobbs S.
30 or Murry L.E. (1992) in McGraw Hill Yearbook of Science and Technology,

McGraw Hill, New York, N.Y., pp 191-196; or Weissbach and Weissbach (1988) *Methods for Plant Molecular Biology*, Academic Press, New York, N.Y., pp 421-463.

Variant product may also be expressed in an insect system. In one such system, *Autographa californica* nuclear polyhedrosis virus (AcNPV) is used as a vector to express foreign genes in *Spodoptera frugiperda* cells or in *Trichoplusia* larvae. The variant product coding sequence may be cloned into a nonessential region of the virus, such as the polyhedrin gene, and placed under control of the polyhedrin promoter. Successful insertion of variant coding sequence will render the polyhedrin gene inactive and produce recombinant virus lacking coat protein coat. The recombinant viruses are then used to infect *S. frugiperda* cells or *Trichoplusia* larvae in which variant protein is expressed (Smith *et al.*, *J. Virol.* 46:584, (1983); Engelhard, E.K. *et al.*, *Proc. Nat. Acad. Sci.* 91:3224-7, (1994)).

In mammalian host cells, a number of viral-based expression systems may be utilized. In cases where an adenovirus is used as an expression vector, a variant product coding sequence may be ligated into an adenovirus transcription/translation complex consisting of the late promoter and tripartite leader sequence. Insertion in a nonessential E1 or E3 region of the viral genome will result in a viable virus capable of expressing variant protein in infected host cells (Logan and Shenk, *Proc. Natl. Acad. Sci.* 81:3655-59, (1984). In addition, transcription enhancers, such as the Rous sarcoma virus (RSV) enhancer, may be used to increase expression in mammalian host cells.

Specific initiation signals may also be required for efficient translation of a variant product coding sequence. These signals include the ATG initiation codon and adjacent sequences. In cases where variant product coding sequence, its initiation codon and upstream sequences are inserted into the appropriate expression vector, no additional translational control signals may be needed. However, in cases where only coding sequence, or a portion thereof, is inserted, exogenous transcriptional control signals including the ATG initiation codon must be provided. Furthermore, the initiation codon must be in the correct

reading frame to ensure transcription of the entire insert. Exogenous transcriptional elements and initiation codons can be of various origins, both natural and synthetic. The efficiency of expression may be enhanced by the inclusion of enhancers appropriate to the cell system in use (Scharf, D. *et al.*,
5 (1994) *Results Probl. Cell Differ.*, 20:125-62, (1994); Bittner *et al.*, *Methods in Enzymol* 153:516-544, (1987)).

In a further embodiment, the present invention relates to host cells containing the above-described constructs. The host cell can be a higher eukaryotic cell, such as a mammalian cell, or a lower eukaryotic cell, such as a
10 yeast cell, or the host cell can be a prokaryotic cell, such as a bacterial cell. Introduction of the construct into the host cell can be effected by calcium phosphate transfection, DEAE-Dextran mediated transfection, or electroporation (Davis, L., Dibner, M., and Battey, I. (1986) *Basic Methods in Molecular Biology*). Cell-free translation systems can also be employed to produce
15 polypeptides using RNAs derived from the DNA constructs of the present invention.

A host cell strain may be chosen for its ability to modulate the expression of the inserted sequences or to process the expressed protein in the desired fashion. Such modifications of the protein include, but are not limited to,
20 acetylation, carboxylation, glycosylation, phosphorylation, lipidation and acylation. Post-translational processing which cleaves a "pre-pro" form of the protein may also be important for correct insertion, folding and/or function. Different host cells such as CHO, HeLa, MDCK, 293, WI38, etc. have specific cellular machinery and characteristic mechanisms for such post-translational
25 activities and may be chosen to ensure the correct modification and processing of the introduced, foreign protein.

For long-term, high-yield production of recombinant proteins, stable expression is preferred. For example, cell lines which stably express variant product may be transformed using expression vectors which contain viral origins
30 of replication or endogenous expression elements and a selectable marker gene.

Following the introduction of the vector, cells may be allowed to grow for 1-2 days in an enriched media before they are switched to selective media. The purpose of the selectable marker is to confer resistance to selection, and its presence allows growth and recovery of cells which successfully express the introduced sequences. Resistant clumps of stably transformed cells can be proliferated using tissue culture techniques appropriate to the cell type.

Any number of selection systems may be used to recover transformed cell lines. These include, but are not limited to, the herpes simplex virus thymidine kinase (Wigler M., *et al.*, *Cell* 11:223-32, (1977)) and adenine phosphoribosyltransferase (Lowy I., *et al.*, *Cell* 22:817-23, (1980)) genes which can be employed in *tk-* or *aprt-* cells, respectively. Also, antimetabolite, antibiotic or herbicide resistance can be used as the basis for selection; for example, *dhfr* which confers resistance to methotrexate (Wigler M., *et al.*, *Proc. Natl. Acad. Sci.* 77:3567-70, (1980)); *npt*, which confers resistance to the aminoglycosides neomycin and G-418 (Colbere-Garapin, F. *et al.*, *J. Mol. Biol.*, 150:1-14, (1981)) and *als* or *pat*, which confer resistance to chlorsulfuron and phosphinotricin acetyltransferase, respectively (Murry, *supra*). Additional selectable genes have been described, for example, *trpB*, which allows cells to utilize indole in place of tryptophan, or *hisD*, which allows cells to utilize histinol in place of histidine (Hartman S.C. and R.C. Mulligan, *Proc. Natl. Acad. Sci.* 85:8047-51, (1988)). The use of visible markers has gained popularity with such markers as anthocyanins, beta-glucuronidase and its substrate, GUS, and luciferase and its substrates, luciferin and ATP, being widely used not only to identify transformants, but also to quantify the amount of transient or stable protein expression attributable to a specific vector system (Rhodes, C.A. *et al.*, *Methods Mol. Biol.*, 55:121-131, (1995)).

Host cells transformed with a nucleotide sequence encoding variant product may be cultured under conditions suitable for the expression and recovery of the encoded protein from cell culture. The product produced by a recombinant cell may be secreted or contained intracellularly depending on the

sequence and/or the vector used. As will be understood by those of skill in the art, expression vectors containing nucleic acid sequences encoding variant product can be designed with signal sequences which direct secretion of variant product through a prokaryotic or eukaryotic cell membrane.

5 The variant product may also be expressed as a recombinant protein with one or more additional polypeptide domains added to facilitate protein purification. Such purification facilitating domains include, but are not limited to, metal chelating peptides such as histidine-tryptophan modules that allow purification on immobilized metals, protein A domains that allow purification on
10 immobilized immunoglobulin, and the domain utilized in the FLAGS extension/affinity purification system (Immunex Corp, Seattle, Wash.). The inclusion of a protease-cleavable polypeptide linker sequence between the purification domain and variant product is useful to facilitate purification. One such expression vector provides for expression of a fusion protein comprising
15 a variant polypeptide fused to a polyhistidine region separated by an enterokinase cleavage site. The histidine residues facilitate purification on IMIAC (immobilized metal ion affinity chromatography, as described in Porath, *et al.*, *Protein Expression and Purification*, 3:263-281, (1992)) while the enterokinase cleavage site provides a means for isolating variant polypeptide from the fusion
20 protein. *pGEX* vectors (Promega, Madison, Wis.) may also be used to express foreign polypeptides as fusion proteins with glutathione S-transferase (GST). In general, such fusion proteins are soluble and can easily be purified from lysed cells by adsorption to ligand-agarose beads (e.g., glutathione-agarose in the case of GST-fusions) followed by elution in the presence of free ligand.

25 Following transformation of a suitable host strain and growth of the host strain to an appropriate cell density, the selected promoter is induced by appropriate means (e.g., temperature shift or chemical induction) and cells are cultured for an additional period. Cells are typically harvested by centrifugation, disrupted by physical or chemical means, and the resulting crude extract retained
30 for further purification. Microbial cells employed in expression of proteins can

be disrupted by any convenient method, including freeze-thaw cycling, sonication, mechanical disruption, or use of cell lysing agents, or other methods, which are well know to those skilled in the art.

The variant products can be recovered and purified from recombinant cell
5 cultures by any of a number of methods well known in the art, including ammonium sulfate or ethanol precipitation, acid extraction, anion or cation exchange chromatography, phosphocellulose chromatography, hydrophobic interaction chromatography, affinity chromatography, hydroxylapatite chromatography, and lectin chromatography. Protein refolding steps can be used,
10 as necessary, in completing configuration of the mature protein. Finally, high performance liquid chromatography (HPLC) can be employed for final purification steps.

C. Diagnostic applications utilizing nucleic acid sequences

15 The nucleic acid sequences of the present invention may be used for a variety of diagnostic purposes. The nucleic acid sequences may be used to detect and quantitate expression of the variant in patient's cells, e.g. biopsied tissues, by detecting the presence of mRNA coding for variant product. Alternatively, the assay may be used to detect soluble variant in the serum or blood. This assay
20 typically involves obtaining total mRNA from the tissue or serum and contacting the mRNA with a nucleic acid probe. The probe is a nucleic acid molecule of at least 20 nucleotides, preferably 20-30 nucleotides, capable of specifically hybridizing with a sequence included within the sequence of a nucleic acid molecule encoding variant product under hybridizing conditions, detecting the
25 presence of mRNA hybridized to the probe, and thereby detecting the expression of variant. This assay can be used to distinguish between absence, presence, and excess expression of variant product and to monitor levels of variant expression during therapeutic intervention. In addition, the assay may be used to compare the levels of the variant of the invention to the levels of the original sequence from

which it has been varied or to levels of other variants, which comparison may have some physiological meaning.

The invention also contemplates the use of the nucleic acid sequences as a diagnostic for diseases resulting from inherited defective variant sequences, or
5 diseases in which the ratio of the amount of the original sequence from which the variant was varied to the novel variants of the invention is altered. These sequences can be detected by comparing the sequences of the defective (i.e., mutant) variant coding region with that of a normal coding region. Association of the sequence coding for mutant variant product with abnormal variant product
10 activity may be verified. In addition, sequences encoding mutant variant products can be inserted into a suitable vector for expression in a functional assay system (e.g., colorimetric assay, complementation experiments in a variant protein deficient strain of HEK293 cells) as yet another means to verify or identify mutations. Once mutant genes have been identified, one can then screen
15 populations of interest for carriers of the mutant gene.

Individuals carrying mutations in the nucleic acid sequence of the present invention may be detected at the DNA level by a variety of techniques. Nucleic acids used for diagnosis may be obtained from a patient's cells, including but not limited to such as from blood, urine, saliva, placenta, tissue biopsy and autopsy
20 material. Genomic DNA may be used directly for detection or may be amplified enzymatically by using PCR (Saiki, *et al.*, *Nature* 324:163-166, (1986)) prior to analysis. RNA or cDNA may also be used for the same purpose. As an example, PCR primers complementary to the nucleic acid of the present invention can be used to identify and analyze mutations in the gene of the present invention.
25 Deletions and insertions can be detected by a change in size of the amplified product in comparison to the normal genotype.

Point mutations can be identified by hybridizing amplified DNA to radiolabeled RNA of the invention or alternatively, radiolabeled antisense DNA sequences of the invention. Sequence changes at specific locations may also be
30 revealed by nuclease protection assays, such as RNase and S1 protection or the

chemical cleavage method (e.g. Cotton, *et al* *Proc. Natl. Acad. Sci. USA*, 85:4397-4401, (1985)), or by differences in melting temperatures. "*Molecular beacons*" (Kostrikis L.G. *et al.*, *Science* 279:1228-1229, (1998)), hairpin-shaped, single-stranded synthetic oligo- nucleotides containing probe sequences which
5 are complementary to the nucleic acid of the present invention, may also be used to detect point mutations or other sequence changes as well as monitor expression levels of variant product. Such diagnostics would be particularly useful for prenatal testing.

Another method for detecting mutations uses two DNA probes which are
10 designed to hybridize to adjacent regions of a target, with abutting bases, where the region of known or suspected mutation(s) is at or near the abutting bases. The two probes may be joined at the abutting bases, e.g., in the presence of a ligase enzyme, but only if both probes are correctly base paired in the region of probe junction. The presence or absence of mutations is then detectable by the
15 presence or absence of ligated probe.

Also suitable for detecting mutations in the variant product coding sequence are oligonucleotide array methods based on sequencing by hybridization (SBH), as described, for example, in U.S. Patent No. 5,547,839. In a typical method, the DNA target analyte is hybridized with an array of
20 oligonucleotides formed on a microchip. The sequence of the target can then be "read" from the pattern of target binding to the array.

D. Gene mapping utilizing nucleic acid sequences

The nucleic acid sequences of the present invention are also valuable for
25 chromosome identification. The sequence is specifically targeted to and can hybridize with a particular location on an individual human chromosome. Moreover, there is a current need for identifying particular sites on the chromosome. Few chromosome marking reagents based on actual sequence data (repeat polymorphisms) are presently available for marking chromosomal
30 location. The mapping of DNAs to chromosomes according to the present

invention is an important first step in correlating those sequences with genes associated with disease.

Briefly, sequences can be mapped to chromosomes by preparing PCR primers (preferably 20-30 bp) from the variant cDNA. Computer analysis of the 3' untranslated region is used to rapidly select primers that do not span more than one exon in the genomic DNA, which would complicate the amplification process. These primers are then used for PCR screening of somatic cell hybrids containing individual human chromosomes. Only those hybrids containing the human gene corresponding to the primer will yield an amplified fragment.

PCR mapping of somatic cell hybrids or using instead radiation hybrids are rapid procedures for assigning a particular DNA to a particular chromosome. Using the present invention with the same oligonucleotide primers, sublocalization can be achieved with panels of fragments from specific chromosomes or pools of large genomic clones in an analogous manner. Other mapping strategies that can similarly be used to map to its chromosome include *in situ* hybridization, prescreening with labeled flow-sorted chromosomes and preselection by hybridization to construct chromosome specific-cDNA libraries.

Fluorescence *in situ* hybridization (FISH) of a cDNA clone to a metaphase chromosomal spread can be used to provide a precise chromosomal location in one step. This technique can be used with cDNA as short as 50 or 60 bases. For a review of this technique, see Verma *et al.*, *Human Chromosomes: a Manual of Basic Techniques*, (1988) Pergamon Press, New York.

Once a sequence has been mapped to a precise chromosomal location, the physical position of the sequence on the chromosome can be correlated with genetic map data. Such data are found, for example, in the OMIM database (Center for Medical Genetics, Johns Hopkins University, Baltimore, MD and National Center for Biotechnology Information, National Library of Medicine, Bethesda, MD). The OMIM gene map presents the cytogenetic map location of disease genes and other expressed genes. The OMIM database provides information on diseases associated with the chromosomal location. Such

associations include the results of linkage analysis mapped to this interval, and the correlation of translocations and other chromosomal aberrations in this area with the advent of polygenic diseases, such as cancer, in general and prostate cancer in particular.

5

E. Therapeutic applications of nucleic acid sequences

Nucleic acid sequences of the invention may also be used for therapeutic purposes. Turning first to the second aspect of the invention (i.e. inhibition of expression of variant), expression of variant product may be modulated through
10 antisense technology, which controls gene expression through hybridization of complementary nucleic acid sequences, i.e. antisense DNA or RNA, to the control, 5' or regulatory regions of the gene encoding variant product. For example, the 5' coding portion of the nucleic acid sequence sequence which codes for the product of the present invention is used to design an antisense
15 oligonucleotide of from about 10 to 40 base pairs in length. Oligonucleotides derived from the transcription start site, e.g. between positions -10 and +10 from the start site, are preferred. An antisense DNA oligonucleotide is designed to be complementary to a region of the nucleic acid sequence involved in transcription (Lee *et al.*, *Nucl. Acids, Res.*, 6:3073, (1979); Cooney *et al.*, *Science* 241:456,
20 (1988); and Dervan *et al.*, *Science* 251:1360, (1991)), thereby preventing transcription and the production of the variant products. An antisense RNA oligonucleotide hybridizes to the mRNA *in vivo* and blocks translation of the mRNA molecule into the variant products (Okano *J. Neurochem.* 56:560, (1991)). The antisense constructs can be delivered to cells by procedures known
25 in the art such that the antisense RNA or DNA may be expressed *in vivo*. The antisense may be antisense mRNA or DNA sequence capable of coding such antisense mRNA. The antisense mRNA or the DNA coding thereof can be complementary to the full sequence of nucleic acid sequences coding for the variant protein or to a fragment of such a sequence which is sufficient to inhibit
30 production of a protein product.

Turning now to the first aspect of the invention, i.e. expression of variant, expression of variant product may be increased by providing coding sequences for coding for said product under the control of suitable control elements ending its expression in the desired host.

5 The nucleic acid sequences of the invention may be employed in combination with a suitable pharmaceutical carrier. Such compositions comprise a therapeutically effective amount of the compound, and a pharmaceutically acceptable carrier or excipient. Such a carrier includes but is not limited to saline, buffered saline, dextrose, water, glycerol, ethanol, and combinations thereof. The
10 formulation should suit the mode of administration.

 The products of the invention as well as any activators and deactivators compounds (see below) which are polypeptides, may also be employed in accordance with the present invention by expression of such polypeptides *in vivo*, which is often referred to as "*gene therapy*." Cells from a patient may be
15 engineered with a nucleic acid sequence (DNA or RNA) encoding a polypeptide *ex vivo*, with the engineered cells then being provided to a patient to be treated with the polypeptide. Such methods are well-known in the art. For example, cells may be engineered by procedures known in the art by use of a retroviral particle containing RNA encoding a polypeptide of the present invention.

20 Similarly, cells may be engineered *in vivo* for expression of a polypeptide *in vivo* by procedures known in the art. As known in the art, a producer cell for producing a retroviral particle containing RNA encoding the polypeptide of the present invention may be administered to a patient for engineering cells *in vivo* and expression of the polypeptide *in vivo*. These and other methods for
25 administering a product of the present invention by such method should be apparent to those skilled in the art from the teachings of the present invention. For example, the expression vehicle for engineering cells may be other than a retrovirus, for example, an adenovirus which may be used to engineer cells *in vivo* after combination with a suitable delivery vehicle.

Retroviruses from which the retroviral plasmid vectors mentioned above may be derived include, but are not limited to, Moloney Murine Leukemia Virus, spleen necrosis virus, retroviruses such as Rous Sarcoma Virus, Harvey Sarcoma Virus, avian leukosis virus, gibbon ape leukemia virus, human immunodeficiency virus, adenovirus, Myeloproliferative Sarcoma Virus, and mammary tumor virus.

The retroviral plasmid vector is employed to transduce packaging cell lines to form producer cell lines. Examples of packaging cells which may be transfected include, but are not limited to, the *PE501*, *PA317*, *psi-2*, *psi-AM*, *PA12*, *T19-14X*, *VT-19-17-H2*, *psi-CRE*, *psi-CRIP*, *GP+E-86*, *GP+envAm12*, and *DAN* cell lines as described in Miller (*Human Gene Therapy*, Vol. 1, pg. 5-14, (1990)). The vector may transduce the packaging cells through any means known in the art. Such means include, but are not limited to, electroporation, the use of liposomes, and CaPO_4 precipitation. In one alternative, the retroviral plasmid vector may be encapsulated into a liposome, or coupled to a lipid, and then administered to a host.

The producer cell line generates infectious retroviral vector particles which include the nucleic acid sequence(s) encoding the polypeptides. Such retroviral vector particles then may be employed, to transduce eukaryotic cells, either *in vitro* or *in vivo*. The transduced eukaryotic cells will express the nucleic acid sequence(s) encoding the polypeptide. Eukaryotic cells which may be transduced include, but are not limited to, embryonic stem cells, embryonic carcinoma cells, as well as hematopoietic stem cells, hepatocytes, fibroblasts, myoblasts, keratinocytes, endothelial cells, and bronchial epithelial cells.

The genes introduced into cells may be placed under the control of inducible promoters, such as the radiation-inducible Egr-1 promoter, (Maceri, H.J., *et al.*, *Cancer Res.*, **56**(19):4311 (1996)), to stimulate variant production or antisense inhibition in response to radiation, eg., radiation therapy for treating tumors.

Example IV. Variant product

The substantially purified variant product of the invention has been defined above as the product coded from the nucleic acid sequence of the invention. Preferably the amino acid sequence is an amino acid sequence having
5 at least 90% identity, preferably 95% identity to any one of the sequences identified as SEQ ID NO: 27 to SEQ ID NO: 52 provided that the amino acid sequence is not identical to that of the original sequence from which it has been varied. The protein or polypeptide may be in mature and/or modified form, also as defined above. Also contemplated are protein fragments having at least 10
10 contiguous amino acid residues, preferably at least 10-20 residues, derived from the variant product, as well as homologues as explained above.

The sequence variations are preferably those that are considered conserved substitutions, as defined above. Thus, for example, a protein with a sequence having at least 90% identity, preferably 95% sequence identity with any of the
15 products identified as SEQ ID NO: 27 to SEQ ID NO: 52, preferably by utilizing conserved substitutions as defined above is also part of the invention, and provided that it is not identical to the original peptide from which it has been varied. In a more specific embodiment, the protein has or contains any one of the sequence identified as SEQ ID NO: 27 to SEQ ID NO: 52. The variant product
20 may be (i) one in which one or more of the amino acid residues in a sequence listed above are substituted with a conserved or non-conserved amino acid residue (preferably a conserved amino acid residue), or (ii) one in which one or more of the amino acid residues includes a substituent group, or (iii) one in which the variant product is fused with another compound, such as a compound to
25 increase the half-life of the protein (for example, polyethylene glycol (PEG)), or a moiety which serves as targeting means to direct the protein to its target tissue or target cell population (such as an antibody), or (iv) one in which additional amino acids are fused to the variant product. Such fragments, variants and derivatives are deemed to be within the scope of those skilled in the art from the teachings
30 herein.

A. Preparation of variant product

Recombinant methods for producing and isolating the variant product, and fragments of the protein are described above.

- 5 In addition to recombinant production, fragments and portions of variant product may be produced by direct peptide synthesis using solid-phase techniques (cf. Stewart *et al.*, (1969) Solid-Phase Peptide Synthesis, WH Freeman Co, San Francisco; Merrifield J., *J. Am. Chem. Soc.*, **85**:2149-2154, (1963)). In vitro peptide synthesis may be performed using manual techniques or by automation.
- 10 Automated synthesis may be achieved, for example, using Applied Biosystems 431A Peptide Synthesizer (Perkin Elmer, Foster City, Calif.) in accordance with the instructions provided by the manufacturer. Fragments of variant product may be chemically synthesized separately and combined using chemical methods to produce the full length molecule.

15

B. Therapeutic uses and compositions utilizing the variant product

- The variant product of the invention is generally useful in treating diseases and disorders which are characterized by a lower than normal level of variant
- 20 expression, and or diseases which can be cured or ameliorated by raising the level of the variant product, even if the level is normal.

- Variant products or fragments may be administered by any of a number of routes and methods designed to provide a consistent and predictable concentration of compound at the target organ or tissue. The product-containing
- 25 compositions may be administered alone or in combination with other agents, such as stabilizing compounds, and/or in combination with other pharmaceutical agents such as drugs or hormones.

- Variant product-containing compositions may be administered by a number of routes including, but not limited to oral, intravenous, intramuscular,
- 30 transdermal, subcutaneous, topical, sublingual, or rectal means as well as by nasal application. variant product-containing compositions may also be administered

via liposomes. Such administration routes and appropriate formulations are generally known to those of skill in the art.

The product can be given via intravenous or intraperitoneal injection. Similarly, the product may be injected to other localized regions of the body. The product may also be administered via nasal insufflation. Enteral administration is also possible. For such administration, the product should be formulated into an appropriate capsule or elixir for oral administration, or into a suppository for rectal administration.

The foregoing exemplary administration modes will likely require that the product be formulated into an appropriate carrier, including ointments, gels, suppositories. Appropriate formulations are well known to persons skilled in the art.

Dosage of the product will vary, depending upon the potency and therapeutic index of the particular polypeptide selected.

A therapeutic composition for use in the treatment method can include the product in a sterile injectable solution, the polypeptide in an oral delivery vehicle, the product in an aerosol suitable for nasal administration, or the product in a nebulized form, all prepared according to well known methods. Such compositions comprise a therapeutically effective amount of the compound, and a pharmaceutically acceptable carrier or excipient. Such a carrier includes but is not limited to saline, buffered saline, dextrose, water, glycerol, ethanol, and combinations thereof. The product of the invention may also be used to modulate endothelial differentiation and proliferation as well as to modulate apoptosis either *ex vivo* or *in vitro*, for example, in cell cultures.

Example V. Screening methods for activators and deactivators (inhibitors)

The present invention also includes an assay for identifying molecules, such as synthetic drugs, antibodies, peptides, or other molecules, which have a modulating effect on the activity of the variant product, e.g. activators or deactivators of the variant product of the present invention. Such an assay

comprises the steps of providing an variant product encoded by the nucleic acid sequences of the present invention, contacting the variant protein with one or more candidate molecules to determine the candidate molecules modulating effect on the activity of the variant product, and selecting from the molecules a candidate's molecule capable of modulating variant product physiological activity.

The variant product, its catalytic or immunogenic fragments or oligopeptides thereof, can be used for screening therapeutic compounds in any of a variety of drug screening techniques. The fragment employed in such a test may be free in solution, affixed to a solid support, borne on a cell membrane or located intracellularly. The formation of binding complexes, between variant product and the agent being tested, may be measured. Alternatively, the activator or deactivator may work by serving as agonist or antagonist, respectively, of the variant receptor, binding entity or target site, and their effect may be determined in connection with any of the above.

Another technique for drug screening which may be used provides for high throughput screening of compounds having suitable binding affinity to the variant product is described in detail by Geysen in PCT Application WO 84/03564, published on Sep. 13, 1984. In summary, large numbers of different small peptide test compounds are synthesized on a solid substrate, such as plastic pins or some other surface. The peptide test compounds are reacted with the full variant product or with fragments of variant product and washed. Bound variant product is then detected by methods well known in the art. Substantially purified variant product can also be coated directly onto plates for use in the aforementioned drug screening techniques. Alternatively, non-neutralizing antibodies can be used to capture the peptide and immobilize it on a solid support.

Antibodies to the variant product, as described in Example VI below, may also be used in screening assays according to methods well known in the art. For example, a "sandwich" assay may be performed, in which an anti-variant

antibody is affixed to a solid surface such as a microtiter plate and variant product is added. Such an assay can be used to capture compounds which bind to the variant product. Alternatively, such an assay may be used to measure the ability of compounds to influence with the binding of variant product to the variant receptor, and then select those compounds which effect the binding.

Example VI. Anti-variant antibodies

A. Synthesis

In still another aspect of the invention, the purified variant product is used to produce anti-variant antibodies which have diagnostic and therapeutic uses related to the activity, distribution, and expression of the variant product.

Antibodies to the variant product may be generated by methods well known in the art. Such antibodies may include, but are not limited to, polyclonal, monoclonal, chimeric, humanized, single chain, Fab fragments and fragments produced by an Fab expression library. Antibodies, i.e., those which inhibit dimer formation, are especially preferred for therapeutic use.

A fragment of the variant product for antibody induction does not require biological activity but have to feature immunological activity; however, the protein fragment or oligopeptide must be antigenic. Peptides used to induce specific antibodies may have an amino acid sequence consisting of at least five amino acids, preferably at least 10 amino acids of the sequences specified in any one of SEQ ID NO: 27 to SEQ ID NO: 52. Preferably they should mimic a portion of the amino acid sequence of the natural protein and may contain the entire amino acid sequence of a small, naturally occurring molecule. Short stretches of variant protein amino acids may be fused with those of another protein such as keyhole limpet hemocyanin and antibody produced against the chimeric molecule. Procedures well known in the art can be used for the production of antibodies to variant product.

For the production of antibodies, various hosts including goats, rabbits, rats, mice, etc may be immunized by injection with variant product or any

portion, fragment or oligopeptide which retains immunogenic properties. Depending on the host species, various adjuvants may be used to increase immunological response. Such adjuvants include but are not limited to Freund's, mineral gels such as aluminum hydroxide, and surface active substances such as
5 lysolecithin, pluronic polyols, polyanions, peptides, oil emulsions, keyhole limpet hemocyanin, and dinitrophenol. BCG (bacilli Calmette-Guerin) and *Corynebacterium parvum* are potentially useful human adjuvants.

Monoclonal antibodies to variant protein may be prepared using any technique which provides for the production of antibody molecules by continuous
10 cell lines in culture. These include but are not limited to the hybridoma technique originally described by Koehler and Milstein (*Nature* 256:495-497, (1975)), the human B-cell hybridoma technique (Kosbor *et al.*, *Immunol. Today* 4:72, (1983); Cote *et al.*, *Proc. Natl. Acad. Sci.* 80:2026-2030, (1983)) and the EBV-hybridoma technique (Cole, *et al.*, *Mol. Cell Biol.* 62:109-120, (1984)).

15 Techniques developed for the production of "chimeric antibodies", the splicing of mouse antibody genes to human antibody genes to obtain a molecule with appropriate antigen specificity and biological activity can also be used (Morrison *et al.*, *Proc. Natl. Acad. Sci.* 81:6851-6855, (1984); Neuberger *et al.*, *Nature* 312:604-608, (1984); Takeda *et al.*, *Nature* 314:452-454, (1985)).
20 Alternatively, techniques described for the production of single chain antibodies (U.S. Pat. No. 4,946,778) can be adapted to produce single-chain antibodies specific for the variant protein.

Antibodies may also be produced by inducing *in vivo* production in the lymphocyte population or by screening recombinant immunoglobulin libraries or
25 panels of highly specific binding reagents as disclosed in Orlandi *et al.* (*Proc. Natl. Acad. Sci.* 86:3833-3837, 1989)), and Winter G and Milstein C., (*Nature* 349:293-299, (1991)).

Antibody fragments which contain specific binding sites for variant protein may also be generated. For example, such fragments include, but are not
30 limited to, the F(ab')₂ fragments which can be produced by pepsin digestion of

the antibody molecule and the Fab fragments which can be generated by reducing the disulfide bridges of the $F(ab')_2$ fragments. Alternatively, Fab expression libraries may be constructed to allow rapid and easy identification of monoclonal Fab fragments with the desired specificity (Huse W.D. *et al.*, *Science* 5 256:1275-1281, (1989)).

B. Diagnostic applications of antibodies

A variety of protocols for competitive binding or immunoradiometric assays using either polyclonal or monoclonal antibodies with established 10 specificities are well known in the art. Such immunoassays typically involve the formation of complexes between the variant product and its specific antibody and the measurement of complex formation. A two-site, monoclonal-based immunoassay utilizing monoclonal antibodies reactive to two noninterfering epitopes on a specific variant product is preferred, but a competitive binding 15 assay may also be employed. These assays are described in Maddox D.E., *et al.*, (*J. Exp. Med.* 158:1211, (1983)).

Antibodies which specifically bind variant product are useful for the diagnosis of conditions or diseases characterized by expression of the novel variant of the invention (where normally it is not expressed) by over or under 20 expression of variant as well as for detection of diseases in which the proportion between the amount of the variants of the invention and the original sequence from which it varied is altered. Alternatively, such antibodies may be used in assays to monitor patients being treated with variant product, its activators, or its deactivators. Diagnostic assays for variant protein include methods utilizing the 25 antibody and a label to detect variant product in human body fluids or extracts of cells or tissues. The products and antibodies of the present invention may be used with or without modification. Frequently, the proteins and antibodies will be labeled by joining them, either covalently or noncovalently, with a reporter molecule. A wide variety of reporter molecules are known in the art.

A variety of protocols for measuring the variant product, using either polyclonal or monoclonal antibodies specific for the respective protein are known in the art. Examples include enzyme-linked immunosorbent assay (ELISA), radioimmunoassay (RIA), and fluorescent activated cell sorting (FACS). As
5 noted above, a two-site, monoclonal-based immunoassay utilizing monoclonal antibodies reactive to two non-interfering epitopes on variant product is preferred, but a competitive binding assay may be employed. These assays are described, among other places, in Maddox, *et al.* (*supra*). Such protocols provide a basis for diagnosing altered or abnormal levels of variant product expression.

10 Normal or standard values for variant product expression are established by combining body fluids or cell extracts taken from normal subjects, preferably human, with antibody to variant product under conditions suitable for complex formation which are well known in the art. The amount of standard complex formation may be quantified by various methods, preferably by photometric
15 methods. Then, standard values obtained from normal samples may be compared with values obtained from samples from subjects potentially affected by disease. Deviation between standard and subject values establishes the presence of disease state.

The antibody assays are useful to determine the level of variant product
20 present in a body fluid sample, in order to determine whether it is being expressed at all, whether it is being overexpressed or underexpressed in the tissue, or as an indication of how variant levels of variable products are responding to drug treatment.

By another aspect the invention concerns methods for determining the
25 presence or level of various anti-variant antibodies in a biological sample obtained from patients, such as blood or serum sample using as an antigen the variant product. Determination of said antibodies may be indicative to a plurality of pathological conditions or diseases.

C. Therapeutic uses of antibodies

In addition to their diagnostic use the antibodies may have a therapeutical utility in blocking or decreasing the activity of the variant product in pathological conditions where beneficial effect can be achieved by such a decrease.

5 The antibody employed is preferably a humanized monoclonal antibody, or a human Mab produced by known globulin-gene library methods. The antibody is administered typically as a sterile solution by IV injection, although other parenteral routes may be suitable. Typically, the antibody is administered in an amount between about 1-15 mg/kg body weight of the subject. Treatment is
10 continued, e.g., with dosing every 1-7 days, until a therapeutic improvement is seen.

Although the invention has been described with reference to specific methods and embodiments, it is appreciated that various modifications and changes may be made without departing from the invention.

CLAIMS:

1. An isolated nucleic acid sequence, of an alternative splicing variant, selected from the group consisting of:
 - (i) the nucleic acid sequence depicted in any one of SEQ ID NO: 1 to
5 SEQ ID NO: 26;
 - (ii) nucleic acid sequences having at least 90% identity with the sequence of (i) with the proviso that each sequence is different than the original nucleic acid sequence from which the sequences of (i) have been varied by alternative splicing; and
 - 10 (iii) fragments of (i) or (ii) of at least 20 b.p., provided that said fragment contains a sequence which is not present, as a continuous stretch of nucleotides, in the original nucleic acid sequence from which the sequences of (i) have been varied by alternative splicing.
2. An isolated nucleic acid sequence according to Claim 1, having at least 95%
15 identity to any one of SEQ ID NO: 1 to SEQ ID NO: 26.
3. An isolated nucleic acid sequence complementary to the nucleic acid sequence of Claim 1.
4. An amino acid sequence selected from the group consisting of:
 - (i) an amino acid sequence coded by the isolated nucleic acid sequence
20 of alternative splice variants of Claim 1;
 - (ii) homologues of the amino acid sequences of (i) in which one or more amino acids has been added, deleted, replaced or chemically modified in the region or adjacent to the region where the amino acid sequences differs from the original amino acid sequence, coded
25 by the original nucleic acid sequence from which the variant has been varied.
5. An amino acid sequence according to Claim 5, as depicted in any one of SEQ ID NO: 27 to SEQ ID NO: 52.
6. An isolated nucleic acid sequence coding for any one of the amino acid
30 sequences of Claim 5.

7. A purified antibody which binds specifically to any of the amino acid sequence of Claim 4.
8. An expression vector comprising any one of the nucleic acid sequences of Claim 1 and control elements for the expression of the nucleic acid sequence in a suitable host.
9. An expression vector comprising any one of the nucleic acid sequences of Claim 3, and control elements for the expression of the nucleic acid sequences in a suitable host.
10. A host cell transfected by the expression vector of Claim 8.
- 10 11. A host cell transfected by the expression vector of Claim 9.
12. A pharmaceutical composition comprising a pharmaceutically acceptable carrier and as an active ingredient an agent selected from the group consisting of:
 - (i) the expression vector of Claim 8; and
 - (ii) any one of the amino acid sequences of Claim 4.
- 15 13. A pharmaceutical composition according to Claim 12, for treatment of diseases which can be ameliorated or cured by raising the level of any one of the amino acid sequences depicted in SEQ ID NO: 27 to SEQ ID NO: 52.
14. A pharmaceutical composition comprising a pharmaceutically acceptable carrier and as an active ingredient an agent selected from the group consisting of:
 - 20 (i) any one of the nucleic acid sequences of Claim 3;
 - (ii) the expression vector of Claim 9; and
 - (iii) the purified antibody of Claim 7.
15. A pharmaceutical composition according to Claim 14, for treatment of diseases which can be ameliorated or cured by decreasing the level of any one of the amino acid sequences depicted in SEQ ID NO: 27 to SEQ ID NO: 52.
- 25 16. A method for detecting an variant nucleic acid sequence in a biological sample, comprising the steps of:
 - (a) hybridizing to nucleic acid material of said biological sample any one of the nucleic acid sequences of Claim 1; and
 - 30 (b) detecting said hybridization complex;

wherein the presence of said hybridization complex correlates with the presence of an variant nucleic acid sequence in the said biological sample.

17. A method for detecting an variant nucleic acid sequence in a biological sample, comprising the steps of:

- 5 (a) hybridizing to nucleic acid material of said biological sample any one of the nucleic acid sequences of Claim 3; and

(b) detecting said hybridization complex;

wherein the presence of said hybridization complex correlates with the presence of an variant nucleic acid sequence in the said biological sample.

10 18. A method for determining the level of variant nucleic acid sequences in a biological sample comprising the steps of:

(a) hybridizing to nucleic acid material of said biological sample any one of the nucleic acid sequences of Claim 1; and

(b) determining the amount of hybridization complexes and normalizing
15 said amount to provide the level of the variant nucleic acid sequences in the sample.

19. A method for determining the level of variant nucleic acid sequences in a biological sample comprising the steps of:

(a) hybridizing to nucleic acid material of said biological sample any one
20 of the nucleic acid sequences of Claim 3; and

(b) determining the amount of hybridization complexes and normalizing said amount to provide the level of the variant nucleic acid sequences in the sample.

20. A method for determining the ratio between the level of variant of the
25 nucleic acid sequence in a first biological sample and the level of the original sequence from which the variant has been varied by alternative splicing in a second biological sample comprising:

- (a) determining the level of the variant nucleic acid sequence in the first biological sample according to the method of Claim 19;
30 (b) determining the level of the original sequence in the second biological sample; and

(c) comprising the levels obtained in (a) and (b) to give said ratio.

21. A method according to Claim 20, wherein said first and said second biological samples are the same sample.

22. A method according to Claim 20, wherein the nucleic acid material of said
5 biological sample are mRNA transcripts.

23. A method according to Claim 22, where the nucleic acid sequence is present in a nucleic acid chip.

24. A method for identifying candidate compounds capable of binding to the variant product and modulating its activity the method comprising:

- 10 (i) providing any one of the amino acid sequences as defined in Claim 4;
(ii) contacting a candidate compound with said amino acid sequence;
(iii) determining the effect of said candidate compound on the biological activity of said protein or polypeptide and selecting those compounds which show a significant effect on said biological activity.

15 25. A method according to Claim 24, wherein the compound is an activator and the measured effect is increase in the biological activity.

26. A method according to Claim 24, wherein the compound is an deactivator and the effect is decrease in the biological activity.

27. An activator of any one of the amino acid sequences of Claim 4.

20 28. An deactivator of any one of the amino acid sequences of Claim 4.

29. A method for detecting any one of the amino acid sequences of Claim 4 in a biological sample, comprising the steps of:

(a) contacting with said biological sample the antibody of Claim 7, thereby forming an antibody-antigen complex; and

25 (b) detecting said antibody-antigen complex
wherein the presence of said antibody-antigen complex correlates with the presence of the desired amino acid in said biological sample.

30. A method for detecting the level of any one of the amino acid sequence of Claim 4 in a biological sample, comprising the steps of:

30 (a) contacting with said biological sample the antibody of Claim 7, thereby forming an antibody-antigen complex; and

(b) detecting the amount of said antibody-antigen complex and normalizing said amount to provide the level of said amino acid sequence in the sample.

31. A method for determining the ratio between the level of any one of the
5 amino acid sequences of Claim 4 present in a first biological sample and the level of the original amino acid sequences from which they were varied by alternative splicing, present in a second biological sample, the method comprising:

(a) determining the level of the amino acid sequences of Claim 4 into a first sample by the method of Claim 30;

10 (b) determining the level of the original amino acid sequence in the second sample; and

(c) comparing the level obtained in (a) and (b) to give said ratio.

32. A method according to Claim 31, wherein said first and said second biological samples are the same sample.

15 33. A method for detecting any one of the antibodies of Claim 7 in a biological sample comprising the steps of:

(a) contacting said biological sample with any one of the amino acid sequences of Claim 4 thereby forming an antibody-antigen complex; and

(b) detecting said antibody-antigen complex

20 wherein the presence of said antibody-antigen complex correlates with the presence of the antibody in said biological sample.

34. A method for detecting the level of any one of the antibodies of Claim 7 in a biological sample comprising the steps of:

(a) contacting said biological sample with any one of the amino acid
25 sequences of Claim 4;

(b) detecting the amount of said antibody-antigen complex and normalizing said amount to provide the levels of said antibody in the sample.

ABSTRACT

The present invention concerns novel variants produced by alternative splicing of known genes, amino acid sequences of the variants as well as their uses in detection and therapy.

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<210> 5
<211> 556
<212> DNA
<213> Homo sapiens

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<400> 5
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<210> 6
<211> 934
<212> DNA
<213> Homo sapiens

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<400> 6

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<210> 7

<211> 783

<212> DNA

<213> Homo sapiens

<400> 7

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<210> 8

<211> 895

<212> DNA

<213> Homo sapiens

<400> 8

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```

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<210> 9
<211> 2565
<212> DNA
<213> Homo sapiens

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<400> 9
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<210> 10
<211> 2438
<212> DNA
<213> Homo sapiens

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<400> 10

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<210> 11
<211> 2522
<212> DNA
<213> Homo sapiens

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<400> 11
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aa 2522

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<210> 12
<211> 2272
<212> DNA
<213> Homo sapiens

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<400> 12
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<211> 1153

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<210> 18

<211> 3239

<212> DNA

<213> Homo sapiens

<400> 18

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<210> 13
<211> 917
<212> DNA
<213> Homo sapiens

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<400> 19
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<210> 20
<211> 1819
<212> DNA
<213> Homo sapiens

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<400> 20
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<210> 21
<211> 2294
<212> DNA
<213> Homo sapiens

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<210> 22
<211> 594
<212> DNA
<213> Homo sapiens

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<400> 22

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<210> 23

<211> 881

<212> DNA

<213> Homo sapiens

<400> 23

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<210> 24

<211> 893

<212> DNA

<213> Homo sapiens

<400> 24

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<210> 25

<211> 887
 <212> DNA
 <213> Homo sapiens

<400> 25

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<210> 26
 <211> 814
 <212> DNA
 <213> Homo sapiens

<400> 26

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ctagcgctca ggccttgcg ccctcgcgcg agatccagcg cccagagaga caccagatcc 180
tgcaattccg acctcgtcat caggggccag tctgtgggga caccagaagt caaccagacc 240
accttatacc agcgttatga gatcaagatg accaagatgt ataaagggtt ccaagcotta 300
ggggatgcgg ctgacatccg gttcgtctac acccccgcca tggagagtgt ctgaggatcc 360
ttccacaggt cccacacccg cagcgaggag tttctcattg ctggaaaact gcaggatgga 420
ctcttgacac tcaactacct cagtttctgt gctccctgga acagcctgag cttagctcag 480
cgccgggggt tcaccaagac ctacactgtt ggctgtgagg aatgcacagt gtttccctgt 540
ttatccatcc cctgcaaaact gcagagtggc actcattgt tgtggacgga ccagctcctc 600
caaggctctg aaaagggtt ccagtcctgt cacttgcct gctgcctcg ggagccaggg 660
ctgtgcacct ggcagtccct gcggtcccag atagcctgaa tcttgcccgg agtggagact 720
gaagcctgca cagtgtccac cctgttccca ctcccatctt tcttccggac aatgaaataa 780
agagttacca cccagcagaa aaaacaaaca agtc 814

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<210> 27
 <211> 481
 <212> PRT
 <213> Homo sapiens

<400> 27

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Met Arg Glu Asn Met Ala Arg Gly Pro Cys Asn Ala Pro Arg Trp Val
  1             5             10             15

Ser Leu Met Val Leu Val Ala Ile Gly Thr Ala Val Thr Ala Ala Val
      20             25             30

Asn Pro Gly Val Val Val Arg Ile Ser Gln Lys Gly Leu Asp Tyr Ala
      35             40             45

Ser Gln Gln Gly Thr Ala Ala Leu Gln Lys Glu Leu Lys Arg Ile Lys
      50             55             60

```

Ile Pro Asp Tyr Ser Asp Ser Phe Lys Ile Lys His Leu Gly Lys Gly
 65 70 75 80
 His Tyr Ser Phe Tyr Ser Met Asp Ile Arg Glu Phe Gln Leu Pro Ser
 85 90 95
 Ser Gln Ile Ser Met Val Pro Asn Val Gly Leu Lys Phe Ser Ile Ser
 100 105 110
 Asn Ala Asn Ile Lys Ile Ser Gly Lys Trp Lys Ala Gln Lys Arg Phe
 115 120 125
 Leu Lys Met Ser Gly Asn Phe Asp Leu Ser Ile Glu Gly Met Ser Ile
 130 135 140
 Ser Ala Asp Leu Lys Leu Gly Ser Asn Pro Thr Ser Gly Lys Pro Thr
 145 150 155 160
 Ile Thr Cys Ser Ser Cys Ser Ser His Ile Asn Ser Val His Val His
 165 170 175
 Ile Ser Lys Ser Lys Val Gly Trp Leu Ile Gln Leu Phe His Lys Lys
 180 185 190
 Ile Glu Ser Ala Leu Arg Asn Lys Met Asn Ser Gln Val Cys Glu Lys
 195 200 205
 Val Thr Asn Ser Val Ser Ser Lys Leu Gln Pro Tyr Phe Gln Thr Leu
 210 215 220
 Pro Val Met Thr Lys Ile Asp Ser Val Ala Gly Ile Asn Tyr Gly Leu
 225 230 235 240
 Val Ala Pro Pro Ala Thr Thr Ala Glu Thr Leu Asp Val Gln Met Lys
 245 250 255
 Gly Glu Phe Tyr Ser Glu Asn His His Asn Pro Pro Pro Phe Ala Pro
 260 265 270
 Pro Val Met Glu Phe Pro Ala Ala His Asp Arg Met Val Tyr Leu Gly
 275 280 285
 Leu Ser Asp Tyr Phe Phe Asn Thr Ala Gly Leu Val Tyr Gln Glu Ala
 290 295 300
 Gly Val Leu Lys Met Thr Leu Arg Asp Asp Met Ile Pro Lys Glu Ser
 305 310 315 320
 Lys Phe Arg Leu Thr Thr Lys Phe Phe Gly Thr Phe Leu Pro Glu Val
 325 330 335
 Ala Lys Lys Phe Pro Asn Met Lys Ile Gln Ile His Val Ser Ala Ser
 340 345 350
 Thr Pro Pro His Leu Ser Val Gln Pro Thr Gly Leu Thr Phe Tyr Pro
 355 360 365
 Ala Val Asp Val Gln Ala Phe Ala Val Leu Pro Asn Ser Ser Leu Ala
 370 375 380

Ser Leu Phe Leu Ile Gly Met Gly Lys Gln Phe Leu Gly Trp Thr Asp
 385 390 395 400

Glu Glu Pro Gln Thr Val Pro Thr Ala Leu Ser Leu Glu Ser Gly Asp
 405 410 415

His Val Asn Pro Val Trp Ile Gln Thr Trp Thr Val Ser Leu Arg Ser
 420 425 430

Leu Arg Leu Glu Ser Leu Tyr Ser Met Val Pro Thr Pro Gly Gly Ile
 435 440 445

His Ser Pro Ser His Ser Leu Val Arg Leu Phe Thr Tyr Ser Phe Asn
 450 455 460

Tyr Ser Phe Ser Gln Phe Leu Ile His Ser Xaa Ile His Ser Met Leu
 465 470 475 480

Ala

<210> 28
 <211> 628
 <212> PRT
 <213> Homo sapiens

<400> 28
 Met Gly Arg Leu Gln Leu Val Val Leu Gly Leu Thr Cys Cys Trp Ala
 1 5 10 15

Val Ala Ser Ala Ala Lys Leu Gly Ala Val Tyr Thr Glu Gly Gly Phe
 20 25 30

Val Glu Gly Val Asn Lys Lys Leu Gly Leu Leu Gly Asp Ser Val Asp
 35 40 45

Ile Phe Lys Gly Ile Pro Phe Ala Ala Pro Thr Lys Ala Leu Glu Asn
 50 55 60

Pro Gln Pro His Pro Gly Trp Gln Gly Thr Leu Lys Ala Lys Asn Phe
 65 70 75 80

Lys Lys Arg Cys Leu Gln Ala Thr Ile Thr Gln Asp Ser Thr Tyr Gly
 85 90 95

Asp Glu Asp Cys Leu Tyr Leu Asn Ile Trp Val Pro Gln Gly Arg Lys
 100 105 110

Gln Val Ser Arg Asp Leu Pro Val Met Ile Trp Ile Tyr Gly Gly Ala
 115 120 125

Phe Leu Met Gly Ser Gly His Gly Ala Asn Phe Leu Asn Asn Tyr Leu
 130 135 140

Tyr Asp Gly Glu Glu Ile Ala Thr Arg Gly Asn Val Ile Val Val Thr
 145 150 155 160

Phe Asn Tyr Arg Val Gly Pro Leu Gly Phe Leu Ser Thr Gly Asp Ala
 165 170 175

Asn Leu Pro Gly Asn Tyr Gly Leu Arg Asp Gln His Met Ala Ile Ala
 180 185 190
 Trp Val Lys Arg Asn Ile Ala Ala Phe Gly Gly Asp Pro Asn Asn Ile
 195 200 205
 Thr Leu Phe Gly Glu Ser Ala Gly Gly Ala Ser Val Ser Leu Gln Thr
 210 215 220
 Leu Ser Pro Tyr Asn Lys Gly Leu Ile Arg Arg Ala Ile Ser Gln Ser
 225 230 235 240
 Gly Val Ala Leu Ser Pro Trp Val Ile Gln Lys Asn Pro Leu Phe Trp
 245 250 255
 Ala Lys Lys Val Ala Glu Lys Val Gly Cys Pro Val Gly Asp Ala Ala
 260 265 270
 Arg Met Ala Gln Cys Leu Lys Val Thr Asp Pro Arg Ala Leu Thr Leu
 275 280 285
 Ala Tyr Lys Val Pro Leu Ala Gly Leu Glu Tyr Pro Met Leu His Tyr
 290 295 300
 Val Gly Phe Val Pro Val Ile Asp Gly Asp Phe Ile Pro Ala Asp Pro
 305 310 315 320
 Ile Asn Leu Tyr Ala Asn Ala Ala Asp Ile Asp Tyr Ile Ala Gly Thr
 325 330 335
 Asn Asn Met Asp Gly His Ile Phe Ala Ser Ile Asp Met Pro Ala Ile
 340 345 350
 Asn Lys Gly Asn Lys Lys Val Thr Glu Glu Asp Phe Tyr Lys Leu Val
 355 360 365
 Ser Glu Phe Thr Ile Thr Lys Gly Leu Arg Gly Ala Lys Thr Thr Phe
 370 375 380
 Asp Val Tyr Thr Glu Ser Trp Ala Gln Asp Pro Ser Gln Glu Asn Lys
 385 390 395 400
 Lys Lys Thr Val Val Asp Phe Glu Thr Asp Val Leu Phe Leu Val Pro
 405 410 415
 Thr Glu Ile Ala Leu Ala Gln His Arg Ala Asn Ala Lys Ser Ala Lys
 420 425 430
 Thr Tyr Ala Tyr Leu Phe Ser His Pro Ser Arg Met Pro Val Tyr Pro
 435 440 445
 Lys Trp Val Gly Ala Asp His Ala Asp Asp Ile Gln Tyr Val Phe Gly
 450 455 460
 Lys Pro Phe Ala Thr Pro Thr Gly Tyr Arg Pro Gln Asp Arg Thr Val
 465 470 475 480
 Ser Lys Ala Met Ile Ala Tyr Trp Thr Asn Phe Ala Lys Thr Gly Asp
 485 490 495
 Pro Asn Met Gly Asp Ser Ala Val Pro Thr His Trp Glu Pro Tyr Thr

500 505 510
 Thr Glu Asn Ser Gly Tyr Leu Glu Ile Thr Lys Lys Met Gly Ser Ser
 515 520 525
 Ser Met Lys Arg Ser Leu Arg Thr Asn Phe Leu Arg Tyr Trp Thr Leu
 530 535 540
 Thr Tyr Leu Ala Leu Pro Thr Val Thr Asp Gln Glu Ala Thr Pro Val
 545 550 555 560
 Pro Pro Thr Gly Asp Ser Glu Ala Thr Pro Val Pro Pro Thr Gly Asp
 565 570 575
 Ser Glu Thr Ala Pro Val Pro Pro Thr Gly Asp Ser Gly Ala Pro Pro
 580 585 590
 Val Pro Pro Thr Gly Asp Ser Gly Ala Pro Pro Val Pro Pro Thr Gly
 595 600 605
 Asp Ser Gly Ala Pro Pro Val Pro Pro Thr Gly Asp Ser Glu Ala Arg
 610 615 620
 Ala His Leu Gly
 625

 <210> 29
 <211> 641
 <212> PRT
 <213> Homo sapiens

 <400> 29
 Leu Leu Leu Leu Gly Phe Leu Leu Val Ser Leu Glu Ser Thr Leu Ser
 1 5 10 15
 Ile Pro Pro Trp Glu Ala Pro Lys Glu His Lys Tyr Lys Ala Glu Glu
 20 25 30
 His Thr Val Val Leu Thr Val Thr Gly Glu Pro Cys His Phe Pro Phe
 35 40 45
 Gln Tyr His Arg Gln Leu Tyr His Lys Cys Thr His Lys Gly Arg Pro
 50 55 60
 Gly Pro Gln Pro Trp Cys Ala Thr Thr Pro Asn Phe Asp Gln Asp Gln
 65 70 75 80
 Arg Trp Gly Tyr Cys Leu Glu Pro Lys Lys Val Lys Asp His Cys Ser
 85 90 95
 Lys His Ser Pro Cys Gln Lys Gly Gly Thr Cys Val Asn Met Pro Ser
 100 105 110
 Gly Pro His Cys Leu Cys Pro Gln His Leu Thr Gly Asn His Cys Gln
 115 120 125
 Lys Glu Lys Cys Phe Glu Pro Gln Leu Leu Arg Phe Phe His Lys Asn
 130 135 140
 Glu Ile Trp Tyr Arg Thr Glu Gln Ala Ala Val Ala Arg Cys Gln Cys

145	150	155	160
Lys Gly Pro Asp Ala His Cys Gln Arg Leu Ala Ser Gln Ala Cys Arg	165	170	175
Thr Asn Pro Cys Leu His Gly Gly Arg Cys Leu Glu Val Glu Gly His	180	185	190
Arg Leu Cys His Cys Pro Val Gly Tyr Thr Gly Pro Phe Cys Asp Val	195	200	205
Asp Thr Lys Ala Ser Cys Tyr Asp Gly Arg Gly Leu Ser Tyr Arg Gly	210	215	220
Leu Ala Arg Thr Thr Leu Ser Gly Ala Pro Cys Gln Pro Trp Ala Ser	225	230	235
Glu Ala Thr Tyr Arg Asn Val Thr Ala Glu Gln Ala Arg Asn Trp Gly	245	250	255
Leu Gly Gly His Ala Phe Cys Arg Asn Pro Asp Asn Asp Ile Arg Pro	260	265	270
Trp Cys Phe Val Leu Asn Arg Asp Arg Leu Ser Trp Glu Tyr Cys Asp	275	280	285
Leu Ala Gln Cys Gln Thr Pro Thr Gln Ala Ala Pro Pro Thr Pro Val	290	295	300
Ser Pro Arg Leu His Val Pro Leu Met Pro Ala Gln Pro Ala Pro Pro	305	310	315
Lys Pro Gln Pro Thr Thr Arg Thr Pro Pro Gln Ser Gln Thr Pro Gly	325	330	335
Ala Leu Pro Ala Lys Arg Glu Gln Pro Pro Ser Leu Thr Arg Asn Gly	340	345	350
Pro Leu Ser Cys Gly Gln Arg Leu Arg Lys Ser Leu Ser Ser Met Thr	355	360	365
Arg Val Val Gly Gly Leu Val Ala Leu Arg Gly Ala His Pro Tyr Ile	370	375	380
Ala Ala Leu Tyr Trp Gly His Ser Phe Cys Ala Gly Ser Leu Ile Ala	385	390	395
Pro Cys Trp Val Leu Thr Ala Ala His Cys Leu Gln Asp Arg Pro Ala	405	410	415
Pro Glu Asp Leu Thr Val Val Leu Gly Gln Glu Arg Arg Asn His Ser	420	425	430
Cys Glu Pro Cys Gln Thr Leu Ala Val Arg Ser Tyr Arg Leu His Glu	435	440	445
Ala Phe Ser Pro Val Ser Tyr Gln His Asp Leu Ala Leu Leu Arg Leu	450	455	460
Gln Glu Asp Ala Asp Gly Ser Cys Ala Leu Leu Ser Pro Tyr Val Gln	465	470	475
			480

Pro Val Cys Leu Pro Ser Gly Ala Ala Arg Pro Ser Glu Thr Thr Leu
 485 490 495
 Cys Gln Val Ala Gly Trp Gly His Gln Phe Glu Gly Ala Glu Glu Tyr
 500 505 510
 Ala Ser Phe Leu Gln Glu Ala Gln Val Pro Phe Leu Ser Leu Glu Arg
 515 520 525
 Cys Ser Ala Pro Asp Val His Gly Ser Ser Ile Leu Pro Gly Met Leu
 530 535 540
 Cys Ala Gly Phe Leu Glu Gly Gly Thr Asp Ala Cys Ala Gly Glu Leu
 545 550 555 560
 Leu Ala Gly Trp Arg Pro Ser Pro Arg Pro Ser Ala Xaa Ser Gln Val
 565 570 575
 His Ser Ala Asp Cys Val Phe Pro Thr Gln Gly Asp Ser Gly Gly Pro
 580 585 590
 Leu Val Cys Glu Asp Gln Ala Ala Glu Arg Arg Leu Thr Leu Gln Gly
 595 600 605
 Ile Ile Ser Trp Gly Ser Gly Cys Gly Asp Arg Asn Lys Pro Gly Val
 610 615 620
 Tyr Thr Asp Val Ala Tyr Tyr Leu Ala Trp Ile Arg Glu His Thr Val
 625 630 635 640
 Ser

<210> 30
 <211> 164
 <212> PRT
 <213> Homo sapiens

<400> 30
 Met Ala Pro Phe Glu Pro Leu Ala Ser Gly Ile Leu Leu Leu Leu Trp
 1 5 10 15
 Leu Ile Ala Pro Ser Arg Ala Cys Thr Cys Val Pro Pro His Pro Gln
 20 25 30
 Thr Ala Phe Cys Asn Ser Asp Leu Val Ile Arg Ala Lys Phe Val Gly
 35 40 45
 Thr Pro Glu Val Asn Gln Thr Thr Leu Tyr Gln Arg Tyr Glu Ile Lys
 50 55 60
 Met Thr Lys Met Tyr Lys Gly Phe Gln Ala Leu Gly Asp Ala Ala Asp
 65 70 75 80
 Ile Arg Phe Val Tyr Thr Pro Ala Met Glu Ser Val Cys Gly Tyr Phe
 85 90 95
 His Arg Ser His Asn Arg Ser Glu Glu Phe Leu Ile Ala Gly Lys Leu
 100 105 110

Gln Asp Gly Leu Leu His Ile Thr Thr Cys Ser Phe Val Ala Pro Trp
 115 120 125
 Asn Ser Leu Ser Leu Ala Gln Arg Arg Gly Phe Thr Lys Thr Tyr Thr
 130 135 140
 Val Gly Cys Glu Glu Cys Thr Val Phe Pro Cys Ser His Ser His Leu
 145 150 155 160
 Ser Ser Gly Gln

<210> 31
 <211> 123
 <212> PRT
 <213> Homo sapiens

<400> 31
 Met Ala Pro Phe Glu Pro Leu Ala Ser Gly Ile Leu Leu Leu Leu Trp
 1 5 10 15
 Leu Ile Ala Pro Ser Arg Ala Cys Thr Cys Val Pro Pro His Pro Gln
 20 25 30
 Thr Ala Phe Cys Asn Ser Asp Leu Val Ile Arg Ala Lys Phe Val Gly
 35 40 45
 Thr Pro Glu Val Asn Gln Thr Thr Leu Tyr Gln Arg Tyr Glu Ile Lys
 50 55 60
 Met Thr Lys Met Tyr Lys Gly Phe Gln Ala Leu Gly Asp Ala Ala Asp
 65 70 75 80
 Ile Arg Phe Val Tyr Thr Pro Ala Met Glu Ser Val Cys Gly Tyr Phe
 85 90 95
 His Arg Ser His Asn Arg Ser Glu Glu Phe Leu Ile Ala Gly Lys Leu
 100 105 110
 Gln Val Val Met Cys Lys Ser Pro Ser Val Val
 115 120

<210> 32
 <211> 211
 <212> PRT
 <213> Homo sapiens

<400> 32
 Met Ala Pro Phe Glu Pro Leu Ala Ser Gly Ile Leu Leu Leu Leu Trp
 1 5 10 15
 Leu Ile Ala Pro Ser Arg Ala Cys Thr Cys Val Pro Pro His Pro Gln
 20 25 30
 Thr Ala Phe Cys Asn Ser Asp Leu Val Ile Arg Ala Lys Phe Val Gly
 35 40 45
 Thr Pro Glu Val Asn Gln Thr Thr Leu Tyr Gln Arg Tyr Glu Ile Lys

50	55	60
Met Thr Lys Met Tyr Lys Gly Phe Gln Ala Leu Gly Asp Ala Ala Asp		
65	70	75 80
Ile Arg Phe Val Tyr Thr Pro Ala Met Glu Ser Val Cys Gly Tyr Phe		
	85	90 95
His Arg Ser His Asn Arg Ser Glu Glu Phe Leu Ile Leu Leu Gly Lys		
	100	105 110
Leu Gln Asp Gly Ile Phe Ala His Ser Leu Thr Cys Ser Phe Cys Trp		
	115	120 125
Val Pro Trp Glu Asn Ser Leu Ser Leu Ala Gln Arg Arg Gly Phe Thr		
	130	135 140
Lys Thr Tyr Thr Val Gly Cys Glu Glu Cys Thr Val Phe Pro Cys Leu		
145	150	155 160
Ser Ile Pro Cys Lys Leu Gln Ser Gly Thr His Cys Leu Trp Thr Asp		
	165	170 175
Gln Leu Leu Gln Gly Ser Glu Lys Gly Phe Gln Ser Arg His Leu Ala		
	180	185 190
Cys Leu Pro Arg Glu Pro Gly Leu Cys Thr Trp Gln Ser Leu Arg Ser		
	195	200 205
Gln Ile Ala		
210		

<210> 33

<211> 160

<212> PRT

<213> Homo sapiens

<400> 33

Met Ala Pro Phe Glu Pro Leu Ala Ser Gly Ile Leu Leu Leu Leu Trp
1 5 10 15

Leu Ile Ala Pro Ser Arg Ala Cys Thr Cys Val Pro Pro His Pro Gln
20 25 30

Thr Ala Phe Cys Asn Ser Asp Leu Val Ile Arg Ala Lys Phe Val Gly
35 40 45

Thr Pro Glu Val Asn Gln Thr Thr Leu Tyr Gln Arg Tyr Glu Ile Lys
50 55 60

Met Thr Lys Met Tyr Lys Gly Phe Gln Ala Leu Gly Asp Ala Ala Asp
65 70 75 80

Ile Arg Phe Val Tyr Thr Pro Ala Met Glu Ser Val Cys Gly Tyr Phe
85 90 95

His Arg Ser His Asn Arg Ser Glu Glu Phe Leu Ile Leu Ser Ile Pro
100 105 110

Cys Lys Leu Gln Ser Gly Thr His Cys Leu Trp Thr Asp Gln Leu Leu

115 120 125
 Gln Gly Ser Glu Lys Gly Phe Gln Ser Arg His Leu Ala Cys Leu Pro
 130 135 140

Arg Glu Pro Gly Leu Cys Thr Trp Gln Ser Leu Arg Ser Gln Ile Ala
 145 150 155 160

<210> 34
 <211> 197
 <212> PRT
 <213> Homo sapiens

<400> 34
 Met Ala Pro Phe Glu Pro Leu Ala Ser Gly Ile Leu Leu Leu Leu Trp
 1 5 10 15
 Leu Ile Ala Pro Ser Arg Ala Cys Thr Cys Val Pro Pro His Pro Gln
 20 25 30
 Thr Ala Phe Cys Asn Ser Asp Leu Val Ile Arg Ala Lys Phe Val Gly
 35 40 45
 Thr Pro Glu Val Asn Gln Thr Thr Leu Tyr Gln Arg Tyr Glu Ile Lys
 50 55 60
 Met Thr Lys Met Tyr Lys Gly Phe Gln Ala Leu Gly Asp Ala Ala Asp
 65 70 75 80
 Ile Arg Phe Val Tyr Thr Pro Ala Met Glu Ser Val Cys Gly Tyr Phe
 85 90 95
 His Arg Ala Gly Lys Leu Gln Asp Gly Leu Leu His Ile Thr Thr Cys
 100 105 110
 Ser Phe Val Ala Pro Trp Asn Ser Leu Ser Leu Ala Gln Arg Arg Gly
 115 120 125
 Phe Thr Lys Thr Tyr Thr Val Gly Cys Glu Glu Cys Thr Val Phe Pro
 130 135 140
 Cys Leu Ser Ile Pro Cys Lys Leu Gln Ser Gly Thr His Cys Leu Trp
 145 150 155 160
 Thr Asp Gln Leu Leu Gln Gly Ser Glu Lys Gly Phe Gln Ser Arg His
 165 170 175
 Leu Ala Cys Leu Pro Arg Glu Pro Gly Leu Cys Thr Trp Gln Ser Leu
 180 185 190
 Arg Ser Gln Ile Ala
 195

<210> 35
 <211> 494
 <212> PRT

<400> 35

Ser Arg Leu Asn Ser Asn Thr Gln Gly Glu Met Lys Phe Glu Val Glu

305	310	315	320
Asn Leu Ile Leu His Lys Asp Tyr Ser Ala Asp Thr Leu Ala His His	325	330	335
Asn Asp Ile Ala Leu Leu Lys Ile Arg Ser Lys Glu Gly Arg Cys Ala	340	345	350
Gln Pro Ser Arg Thr Ile Gln Thr Ile Cys Leu Pro Ser Met Tyr Asn	355	360	365
Asp Pro Gln Phe Gly Thr Ser Cys Glu Ile Thr Gly Phe Gly Lys Glu	370	375	380
Asn Ser Thr Asp Tyr Leu Tyr Pro Glu Gln Leu Lys Met Thr Val Val	385	390	395
Lys Leu Ile Ser His Arg Glu Cys Gln Gln Pro His Tyr Tyr Gly Ser	405	410	415
Glu Val Thr Thr Lys Met Leu Cys Ala Ala Asp Pro Gln Trp Lys Thr	420	425	430
Asp Ser Cys Gln Gly Asp Ser Gly Gly Pro Leu Val Cys Ser Leu Gln	435	440	445
Gly Arg Met Thr Leu Thr Gly Ile Val Ser Trp Gly Arg Gly Cys Ala	450	455	460
Leu Lys Asp Lys Pro Gly Val Tyr Thr Arg Val Ser His Phe Leu Pro	465	470	475
Trp Ile Arg Ser His Thr Lys Glu Glu Asn Gly Leu Ala Leu	485	490	

<210> 36
 <211> 285
 <212> PRT
 <213> Homo sapiens

<400> 36
 Met Gln Met Ser Pro Ala Leu Thr Cys Leu Val Leu Gly Leu Ala Leu
 1 5 10 15
 Val Phe Gly Glu Gly Ser Ala Val His His Pro Pro Ser Tyr Val Ala
 20 25 30
 His Leu Ala Ser Asp Phe Gly Val Arg Val Phe Gln Gln Val Ala Gln
 35 40 45
 Ala Ser Lys Asp Arg Asn Val Val Phe Ser Pro Tyr Gly Val Ala Ser
 50 55 60
 Val Leu Ala Met Leu Gln Leu Thr Thr Gly Gly Glu Thr Gln Gln Gln
 65 70 75 80
 Ile Gln Ala Ala Met Gly Phe Lys Ile Asp Asp Lys Gly Met Ala Pro
 85 90 95
 Ala Leu Arg His Leu Tyr Lys Glu Leu Met Gly Pro Trp Asn Lys Asp

100					105					110						
Glu	Ile	Ser	Thr	Thr	Asp	Ala	Ile	Phe	Val	Gln	Arg	Asp	Leu	Lys	Leu	
115					120					125						
Val	Gln	Gly	Phe	Met	Pro	His	Phe	Phe	Arg	Leu	Phe	Arg	Ser	Thr	Val	
130					135					140						
Lys	Gln	Val	Asp	Phe	Ser	Glu	Val	Glu	Arg	Ala	Arg	Phe	Ile	Ile	Asn	
145					150					155					160	
Asp	Trp	Val	Lys	Thr	His	Thr	Lys	Gly	Met	Ile	Ser	Asn	Leu	Leu	Gly	
165					170					175						
Lys	Gly	Ala	Val	Asp	Gln	Leu	Thr	Arg	Leu	Val	Leu	Val	Asn	Ala	Leu	
180					185					190						
Tyr	Phe	Asn	Gly	Gln	Trp	Lys	Thr	Pro	Phe	Pro	Asp	Ser	Ser	Thr	His	
195					200					205						
Arg	Arg	Leu	Phe	His	Lys	Ser	Asp	Gly	Ser	Thr	Val	Ser	Val	Pro	Met	
210					215					220						
Met	Ala	Gln	Thr	Asn	Lys	Phe	Asn	Tyr	Thr	Glu	Phe	Thr	Thr	Pro	Asp	
225					230					235					240	
Gly	His	Tyr	Tyr	Asp	Ile	Leu	Glu	Leu	Pro	Tyr	His	Gly	Asp	Thr	Leu	
245					250					255						
Ser	Met	Phe	Ile	Ala	Ala	Asp	Leu	Val	Pro	Thr	Glu	Ala	Leu	Cys	Arg	
260					265					270						
Met	Glu	Leu	Arg	Gly	Leu	Gln	Glu	Leu	Leu	Cys	Ala	Trp				
275					280					285						

<210> 37

<211> 399

<212> PRT

<213> Homo sapiens

<400> 37

Met	Gln	Met	Ser	Pro	Ala	Leu	Thr	Cys	Leu	Val	Leu	Gly	Leu	Ala	Leu
1					5				10					15	

Val	Phe	Gly	Glu	Gly	Ser	Ala	Val	His	His	Pro	Pro	Ser	Tyr	Val	Ala
		20						25					30		

His	Leu	Ala	Ser	Asp	Phe	Gly	Val	Arg	Val	Phe	Gln	Gln	Val	Ala	Gln
		35					40					45			

Ala	Ser	Lys	Asp	Arg	Asn	Val	Val	Phe	Ser	Pro	Tyr	Gly	Val	Ala	Ser
		50				55					60				

Val	Leu	Ala	Met	Leu	Cln	Leu	Thr	Thr	Gly	Gly	Glu	Thr	Gln	Gln	Gln
65					70					75					80

Ile	Gln	Ala	Ala	Met	Gly	Phe	Lys	Ile	Asp	Asp	Lys	Gly	Met	Ala	Pro
				85					90					95	

Ala Leu Arg His Leu Tyr Lys Glu Leu Met Gly Pro Trp Asn Lys Asp

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<210> 38
<211> 317
<212> PRT
<213> Homo sapiens
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<400> 38

Met Thr Ala Ala Ser Met Gly Pro Val Arg Val Ala Phe Val Val Leu
 1 5 10 15
 Leu Ala Leu Cys Ser Arg Pro Ala Val Gly Gln Asn Cys Ser Gly Pro
 20 25 30
 Cys Arg Cys Pro Asp Glu Pro Ala Pro Arg Cys Pro Ala Gly Val Ser
 35 40 45
 Leu Val Leu Asp Gly Cys Gly Cys Cys Arg Val Cys Ala Lys Gln Leu
 50 55 60
 Gly Glu Leu Cys Thr Glu Arg Asp Pro Cys Asp Pro His Lys Gly Leu
 65 70 75 80
 Phe Cys Asp Phe Gly Ser Pro Ala Asn Arg Lys Ile Gly Val Cys Thr
 85 90 95
 Ala Lys Asp Gly Ala Pro Cys Ile Phe Gly Gly Thr Val Tyr Arg Ser
 100 105 110
 Gly Glu Ser Phe Gln Ser Ser Cys Lys Tyr Gln Cys Thr Cys Leu Asp
 115 120 125
 Gly Ala Val Gly Cys Met Pro Leu Cys Ser Met Asp Val Arg Leu Pro
 130 135 140
 Ser Pro Asp Cys Pro Leu Pro Leu Glu Asp Thr Phe Gly Pro Asp Pro
 145 150 155 160
 Thr Met Ile Arg Ala Asn Cys Leu Val Gln Thr Thr Glu Trp Ser Ala
 165 170 175
 Cys Ser Lys Thr Cys Gly Met Gly Ile Ser Thr Arg Val Thr Asn Asp
 180 185 190
 Asn Ala Ser Cys Arg Leu Glu Lys Gln Ser Arg Leu Cys Met Val Arg
 195 200 205
 Pro Cys Glu Ser Asp Leu Glu Glu Asn Ile Lys Lys Gly Lys Lys Cys
 210 215 220
 Ile Arg Thr Pro Lys Ile Ser Lys Pro Ile Lys Phe Glu Leu Ser Gly
 225 230 235 240
 Cys Thr Ser Met Lys Thr Tyr Arg Ala Lys Phe Cys Gly Val Cys Thr
 245 250 255
 Asp Gly Arg Cys Cys Thr Pro His Arg Thr Thr Thr Leu Pro Val Glu
 260 265 270
 Phe Lys Cys Pro Asp Gly Glu Val Met Lys Lys Asn Met Met Phe Ile
 275 280 285
 Lys Thr Cys Ala Cys His Tyr Asn Cys Pro Gly Asp Asn Asp Ile Phe
 290 295 300
 Glu Ser Leu Tyr Tyr Arg Lys Met Tyr Gly Asp Met Ala
 305 310 315

<210> 39
 <211> 342
 <212> PRT
 <213> Homo sapiens

<400> 39

Asn	Met	Glu	Asn	Ser	Leu	Arg	Cys	Val	Trp	Val	Pro	Lys	Leu	Ala	Phe	1	5	10	15
Val	Leu	Phe	Gly	Ala	Ser	Leu	Leu	Ser	Ala	His	Leu	Gln	Val	Thr	Gly	20	25	30	
Phe	Gln	Ile	Lys	Ala	Phe	Thr	Ala	Leu	Arg	Phe	Leu	Ser	Glu	Pro	Ser	35	40	45	
Asp	Ala	Val	Thr	Met	Arg	Gly	Gly	Asn	Val	Leu	Leu	Asp	Cys	Ser	Ala	50	55	60	
Glu	Ser	Asp	Arg	Gly	Val	Pro	Val	Ile	Lys	Trp	Lys	Lys	Asp	Ala	Ile	65	70	75	
His	Leu	Ala	Leu	Gly	Met	Asp	Glu	Arg	Lys	Gln	Gln	Leu	Ser	Asn	Gly	85	90	95	
Ser	Leu	Leu	Ile	Gln	Asn	Ile	Leu	His	Ser	Arg	His	His	Lys	Pro	Asp	100	105	110	
Glu	Gly	Leu	Tyr	Gln	Cys	Glu	Ala	Ser	Leu	Gly	Asp	Ser	Gly	Ser	Ile	115	120	125	
Ile	Ser	Arg	Thr	Ala	Lys	Val	Ala	Val	Ala	Gly	Pro	Leu	Arg	Phe	Leu	130	135	140	
Ser	Gln	Thr	Glu	Ser	Val	Thr	Ala	Phe	Met	Gly	Asp	Thr	Val	Leu	Leu	145	150	155	
Lys	Cys	Glu	Val	Ile	Gly	Glu	Pro	Met	Pro	Thr	Ile	His	Trp	Gln	Lys	165	170	175	
Asn	Gln	Gln	Asp	Leu	Thr	Pro	Ile	Pro	Gly	Asp	Ser	Arg	Val	Val	Val	180	185	190	
Leu	Pro	Ser	Gly	Ala	Leu	Gln	Ile	Ser	Arg	Leu	Gln	Pro	Gly	Asp	Ile	195	200	205	
Gly	Ile	Tyr	Arg	Cys	Ser	Ala	Arg	Asn	Pro	Ala	Ser	Ser	Arg	Thr	Gly	210	215	220	
Asn	Glu	Ala	Glu	Val	Arg	Ile	Leu	Ser	Asp	Pro	Gly	Leu	His	Arg	Gln	225	230	235	
Leu	Tyr	Phe	Leu	Gln	Arg	Pro	Ser	Asn	Val	Val	Ala	Ile	Glu	Gly	Lys	245	250	255	
Asp	Ala	Val	Leu	Glu	Cys	Cys	Val	Ser	Gly	Tyr	Pro	Pro	Pro	Ser	Phe	260	265	270	
Thr	Trp	Leu	Arg	Gly	Glu	Glu	Val	Ile	Gln	Leu	Arg	Ser	Lys	Lys	Tyr	275	280	285	

Ser Leu Leu Gly Gly Ser Asn Leu Leu Ile Ser Asn Val Thr Asp Asp
290 295 300

Asp Ser Gly Met Tyr Thr Cys Val Val Thr Tyr Lys Asn Glu Asn Ile
305 310 315 320

Ser Ala Ser Ala Glu Leu Thr Val Leu Val Ile Ile Asp Lys Val Leu
325 330 335

Val Asp Thr Phe Trp Val
340

<210> 40

<211> 1433

<212> PRT

<213> Homo sections

<400> 40

Asn Met Glu Asn Ser Leu Arg Cys Val Trp Val Pro Lys Leu Ala Phe
1 5 10 15

Val Leu Phe Gly Ala Ser Leu Leu Ser Ala His Leu Gln Val Thr Gly
20 25 30

Phe Gln Ile Lys Ala Phe Thr Ala Leu Arg Phe Leu Ser Glu Pro Ser
35 40 45

Asp Ala Val Thr Met Arg Gly Gly Asn Val Leu Leu Asp Cys Ser Ala
50 55 60

Glu Ser Asp Arg Gly Val Pro Val Ile Lys Trp Lys Lys Asp Ala Ile
65 70 75 80

His Leu Ala Leu Gly Met Asp Glu Arg Lys Gln Gln Leu Ser Asn Gly
85 90 95

Ser Leu Leu Ile Gln Asn Ile Leu His Ser Arg His His Lys Pro Asp
100 105 110

Glu Gly Leu Tyr Gln Cys Glu Ala Ser Leu Gly Asp Ser Gly Ser Ile
115 120 125

Ile Ser Arg Thr Ala Lys Val Ala Val Ala Gly Pro Leu Arg Phe Leu
130 135 140

Ser Gln Thr Glu Ser Val Thr Ala Phe Met Gly Asp Thr Val Leu Leu
145 150 155 160

Lys Cys Glu Val Ile Gly Glu Pro Met Pro Thr Ile His Trp Gln Lys
165 170 175

Asn Gln Gln Asp Leu Thr Pro Ile Pro Gly Asp Ser Arg Val Val Val
180 185 190

Leu Pro Ser Gly Ala Leu Gln Ile Ser Arg Leu Gln Pro Gly Asp Ile
195 200 205

Gly Ile Tyr Arg Cys Ser Ala Arg Asn Pro Ala Ser Ser Arg Thr Gly
210 215 220

Asn Glu Ala Glu Val Arg Ile Leu Ser Asp Pro Gly Leu His Arg Gln
 225 230 235 240
 Leu Tyr Phe Leu Gln Arg Pro Ser Asn Val Val Ala Ile Glu Gly Lys
 245 250 255
 Asp Ala Val Leu Glu Cys Cys Val Ser Gly Tyr Pro Pro Pro Ser Phe
 260 265 270
 Thr Trp Leu Arg Gly Glu Glu Val Ile Gln Leu Arg Ser Lys Lys Tyr
 275 280 285
 Ser Leu Leu Gly Gly Ser Asn Leu Leu Ile Ser Asn Val Thr Asp Asp
 290 295 300
 Asp Ser Gly Met Tyr Thr Cys Val Val Thr Tyr Lys Asn Glu Asn Ile
 305 310 315 320
 Ser Ala Ser Ala Glu Leu Thr Val Leu Val Pro Pro Trp Phe Leu Asn
 325 330 335
 His Pro Ser Asn Leu Tyr Ala Tyr Glu Ser Met Asp Ile Glu Phe Glu
 340 345 350
 Cys Thr Val Ser Gly Lys Pro Val Pro Thr Val Asn Trp Met Lys Asn
 355 360 365
 Gly Asp Val Val Ile Pro Ser Asp Tyr Phe Gln Ile Val Gly Gly Ser
 370 375 380
 Asn Leu Arg Ile Leu Gly Val Val Lys Ser Asp Glu Gly Phe Tyr Gln
 385 390 395 400
 Cys Val Ala Glu Asn Glu Ala Gly Asn Ala Gln Thr Ser Ala Gln Leu
 405 410 415
 Ile Val Pro Lys Pro Ala Ile Pro Ser Ser Ser Val Leu Pro Ser Ala
 420 425 430
 Pro Arg Asp Val Val Pro Val Leu Val Ser Ser Arg Phe Val Arg Leu
 435 440 445
 Ser Trp Arg Pro Pro Ala Glu Ala Lys Gly Asn Ile Gln Thr Phe Thr
 450 455 460
 Val Phe Phe Ser Arg Glu Gly Asp Asn Arg Glu Arg Ala Leu Asn Thr
 465 470 475 480
 Thr Gln Pro Gly Ser Leu Glu Leu Thr Val Gly Asn Leu Lys Pro Glu
 485 490 495
 Ala Met Tyr Thr Phe Arg Val Val Ala Tyr Asn Glu Trp Gly Pro Gly
 500 505 510
 Glu Ser Ser Gln Pro Ile Lys Val Ala Thr Gln Pro Glu Leu Gln Val
 515 520 525
 Pro Gly Pro Val Glu Asn Leu Glu Ala Val Ser Thr Ser Pro Thr Ser
 530 535 540

Ile Leu Ile Thr Trp Glu Pro Pro Ala Tyr Ala Asn Gly Pro Val Gln
 545 550 555 560
 Gly Tyr Arg Leu Phe Cys Thr Glu Val Ser Thr Gly Lys Glu Gln Asn
 565 570 575
 Ile Glu Val Asp Gly Leu Ser Tyr Lys Leu Glu Gly Leu Lys Lys Phe
 580 585 590
 Thr Glu Tyr Ser Leu Arg Phe Leu Ala Tyr Asn Arg Tyr Gly Pro Gly
 595 600 605
 Val Ser Thr Asp Asp Ile Thr Val Val Thr Leu Ser Asp Val Pro Ser
 610 615 620
 Ala Pro Pro Gln Asn Val Ser Leu Glu Val Val Asn Ser Arg Ser Ile
 625 630 635 640
 Lys Val Ser Trp Leu Pro Pro Pro Ser Gly Thr Gln Asn Gly Phe Ile
 645 650 655
 Thr Gly Tyr Lys Ile Arg His Arg Lys Thr Thr Arg Arg Gly Glu Met
 660 665 670
 Glu Thr Leu Glu Pro Asn Asn Leu Trp Tyr Leu Phe Thr Gly Leu Glu
 675 680 685
 Lys Gly Ser Gln Tyr Ser Phe Gln Val Ser Ala Met Thr Val Asn Gly
 690 695 700
 Thr Gly Pro Pro Ser Asn Trp Tyr Thr Ala Glu Thr Pro Glu Asn Asp
 705 710 715 720
 Leu Asp Glu Ser Gln Val Pro Asp Gln Pro Ser Ser Leu His Val Arg
 725 730 735
 Pro Gln Thr Asn Cys Ile Ile Met Ser Trp Thr Pro Pro Leu Asn Pro
 740 745 750
 Asn Ile Val Val Arg Gly Tyr Ile Ile Gly Tyr Gly Val Gly Ser Pro
 755 760 765
 Tyr Ala Glu Thr Val Arg Val Asp Ser Lys Gln Arg Tyr Tyr Ser Ile
 770 775 780
 Glu Arg Leu Glu Ser Ser Ser His Tyr Val Ile Ser Leu Lys Ala Phe
 785 790 795 800
 Asn Asn Ala Gly Glu Gly Val Pro Leu Tyr Glu Ser Ala Thr Thr Arg
 805 810 815
 Ser Ile Thr Asp Pro Thr Asp Pro Val Asp Tyr Tyr Pro Leu Leu Asp
 820 825 830
 Asp Phe Pro Thr Ser Val Pro Asp Leu Ser Thr Pro Met Leu Pro Pro
 835 840 845
 Val Gly Val Gln Ala Val Ala Leu Thr His Asp Ala Val Arg Val Ser
 850 855 860
 Trp Ala Asp Asn Ser Val Pro Lys Asn Gln Lys Thr Ser Glu Val Arg

865	870	875	880
Leu Tyr Thr Val Arg Trp Arg Thr Ser Phe Ser Ala Ser Ala Lys Tyr	885	890	895
Lys Ser Glu Asp Thr Thr Ser Leu Ser Tyr Thr Ala Thr Gly Leu Lys	900	905	910
Pro Asn Thr Met Tyr Glu Phe Ser Val Met Val Thr Lys Asn Arg Arg	915	920	925
Ser Ser Thr Trp Ser Met Thr Ala His Ala Thr Thr Tyr Glu Ala Ala	930	935	940
Pro Thr Ser Ala Pro Lys Asp Phe Thr Val Ile Thr Arg Glu Gly Lys	945	950	955
Pro Arg Ala Val Ile Val Ser Trp Gln Pro Pro Leu Glu Ala Asn Gly	965	970	975
Lys Ile Thr Ala Tyr Ile Leu Phe Tyr Thr Leu Asp Lys Asn Ile Pro	980	985	990
Ile Asp Asp Trp Ile Met Glu Thr Ile Ser Gly Asp Arg Leu Thr His	995	1000	1005
Gln Ile Met Asp Leu Asn Leu Asp Thr Met Tyr Tyr Phe Arg Ile Gln	1010	1015	1020
Ala Arg Asn Ser Lys Gly Val Gly Pro Leu Ser Asp Pro Ile Leu Phe	1025	1030	1035
Arg Thr Leu Lys Val Glu His Pro Asp Lys Met Ala Asn Asp Gln Gly	1045	1050	1055
Arg His Gly Asp Gly Gly Tyr Trp Pro Val Asp Thr Asn Leu Ile Asp	1060	1065	1070
Arg Ser Thr Leu Asn Glu Pro Pro Ile Gly Gln Met His Pro Pro His	1075	1080	1085
Gly Ser Val Thr Pro Gln Lys Asn Ser Asn Leu Leu Val Ile Ile Val	1090	1095	1100
Val Thr Val Gly Val Ile Thr Val Leu Val Val Val Ile Val Ala Val	1105	1110	1115
Ile Cys Thr Arg Arg Ser Ser Ala Gln Gln Arg Lys Lys Arg Ala Thr	1125	1130	1135
His Ser Ala Gly Lys Arg Lys Gly Ser Gln Lys Asp Leu Arg Pro Pro	1140	1145	1150
Asp Leu Trp Ile His His Glu Glu Met Glu Met Lys Asn Ile Glu Lys	1155	1160	1165
Pro Ser Gly Thr Asp Pro Ala Gly Arg Asp Ser Pro Ile Gln Ser Cys	1170	1175	1180
Gln Asp Leu Thr Pro Val Ser His Ser Gln Ser Glu Thr Gln Leu Gly	1185	1190	1195
			1200

Ser Lys Ser Thr Ser His Ser Gly Gln Asp Thr Glu Glu Ala Gly Ser
1205 1210 1215

Ser Met Ser Thr Leu Glu Arg Ser Leu Ala Ala Arg Arg Ala Pro Arg
1220 1225 1230

Ala Lys Leu Met Ile Pro Met Asp Ala Gln Ser Asn Asn Pro Ala Val
1235 1240 1245

Val Ser Ala Ile Pro Val Pro Thr Leu Glu Ser Ala Gln Tyr Pro Gly
1250 1255 1260

Ile Leu Pro Ser Pro Thr Cys Gly Tyr Pro His Pro Gln Phe Thr Leu
1265 1270 1275 1280

Arg Pro Val Pro Phe Pro Thr Leu Ser Val Asp Arg Gly Phe Gly Ala
1285 1290 1295

Gly Arg Ser Gln Ser Val Ser Glu Gly Pro Thr Thr Gln Gln Pro Pro
1300 1305 1310

Met Leu Pro Pro Ser Gln Pro Glu His Ser Ser Ser Glu Glu Ala Pro
1315 1320 1325

Ser Arg Thr Ile Pro Thr Ala Cys Val Arg Pro Thr His Pro Leu Arg
1330 1335 1340

Ser Phe Ala Asn Pro Leu Leu Pro Pro Pro Met Ser Ala Ile Glu Pro
1345 1350 1355 1360

Lys Val Pro Tyr Thr Pro Leu Leu Ser Gln Pro Gly Pro Thr Leu Pro
1365 1370 1375

Lys Thr His Val Lys Thr Ala Ser Leu Gly Leu Ala Gly Lys Ala Arg
1380 1385 1390

Ser Pro Leu Leu Pro Val Ser Val Pro Thr Ala Pro Glu Val Ser Glu
1395 1400 1405

Glu Ser His Lys Pro Thr Glu Asp Ser Ala Asn Val Ser Ala Ser Leu
1410 1415 1420

Lys Phe Met Leu His Gln Gly Thr Asp
1425 1430

<210> 41

<211> 865

<212> FRT

<213> Homo sapiens

<400> 41

Met Pro Gly Lys Arg Gly Leu Gly Trp Trp Trp Ala Arg Leu Pro Leu
1 5 10 15

Cys Leu Leu Leu Ser Leu Tyr Gly Pro Trp Met Pro Ser Ser Leu Gly
20 25 30

Lys Pro Lys Gly His Pro His Met Asn Ser Ile Arg Ile Asp Gly Asp
35 40 45

Ile Thr Leu Gly Gly Leu Phe Pro Val His Gly Arg Gly Ser Glu Gly
 50 55 60
 Lys Pro Cys Gly Glu Leu Lys Lys Glu Lys Gly Ile His Arg Leu Glu
 65 70 75 80
 Ala Met Leu Phe Ala Leu Asp Arg Ile Asn Asn Asp Pro Asp Leu Leu
 85 90 95
 Pro Asn Ile Thr Leu Gly Ala Arg Ile Leu Asp Thr Cys Ser Arg Asp
 100 105 110
 Thr His Ala Leu Glu Gln Ser Leu Thr Phe Val Gln Ala Leu Ile Glu
 115 120 125
 Lys Asp Gly Thr Glu Val Arg Cys Gly Ser Gly Gly Pro Pro Ile Ile
 130 135 140
 Thr Lys Pro Glu Arg Val Val Gly Val Ile Gly Ala Ser Gly Ser Ser
 145 150 155 160
 Val Ser Ile Met Val Ala Asn Ile Leu Arg Leu Phe Lys Ile Pro Gln
 165 170 175
 Ile Ser Tyr Ala Ser Thr Ala Pro Asp Leu Ser Asp Asn Ser Arg Tyr
 180 185 190
 Asp Phe Phe Ser Arg Val Val Pro Ser Asp Thr Tyr Gln Ala Gln Ala
 195 200 205
 Met Val Asp Ile Val Arg Ala Leu Lys Trp Asn Tyr Val Ser Thr Val
 210 215 220
 Ala Ser Glu Gly Ser Tyr Gly Glu Ser Gly Val Glu Ala Phe Ile Gln
 225 230 235 240
 Lys Ser Arg Glu Asp Gly Gly Val Cys Ile Ala Gln Ser Val Lys Ile
 245 250 255
 Pro Arg Glu Pro Lys Ala Gly Glu Phe Asp Lys Ile Ile Arg Arg Leu
 260 265 270
 Leu Glu Thr Ser Asn Ala Arg Ala Val Ile Ile Phe Ala Asn Glu Asp
 275 280 285
 Asp Ile Arg Arg Val Leu Glu Ala Ala Arg Arg Ala Asn Gln Thr Gly
 290 295 300
 His Phe Phe Trp Met Gly Ser Asp Ser Trp Gly Ser Lys Ile Ala Pro
 305 310 315 320
 Val Leu His Leu Glu Glu Val Ala Glu Gly Ala Val Thr Ile Leu Pro
 325 330 335
 Lys Arg Met Ser Val Arg Asp Arg Glu Arg Ile Gly Gln Asp Ser Ala
 340 345 350
 Tyr Glu Gln Glu Gly Lys Val Gln Phe Val Ile Asp Ala Val Tyr Ala
 355 360 365

Met Gly His Ala Leu His Ala Met His Arg Asp Leu Cys Pro Gly Arg
 370 375 380
 Val Gly Leu Cys Pro Arg Met Asp Pro Val Asp Gly Thr Gln Leu Leu
 385 390 395 400
 Lys Tyr Ile Arg Asn Val Asn Phe Ser Gly Ile Ala Gly Asn Pro Val
 405 410 415
 Thr Phe Asn Glu Asn Gly Asp Ala Pro Gly Arg Tyr Asp Ile Tyr Gln
 420 425 430
 Tyr Gln Leu Arg Asn Asp Ser Ala Glu Tyr Lys Val Ile Gly Ser Trp
 435 440 445
 Thr Asp His Leu His Leu Arg Ile Glu Arg Met His Trp Pro Gly Ser
 450 455 460
 Gly Gln Gln Leu Pro Arg Ser Ile Cys Ser Leu Pro Cys Gln Pro Gly
 465 470 475 480
 Glu Arg Lys Lys Thr Val Lys Gly Met Pro Cys Cys Trp His Cys Glu
 485 490 495
 Pro Cys Thr Gly Tyr Gln Tyr Gln Val Asp Arg Tyr Thr Cys Lys Thr
 500 505 510
 Cys Pro Tyr Asp Met Arg Pro Thr Glu Asn Arg Thr Gly Cys Arg Pro
 515 520 525
 Ile Pro Ile Ile Lys Leu Glu Trp Gly Ser Pro Trp Ala Val Leu Pro
 530 535 540
 Leu Phe Leu Ala Val Val Gly Ile Ala Ala Thr Leu Phe Val Val Ile
 545 550 555 560
 Thr Phe Val Arg Tyr Asn Asp Thr Pro Ile Val Lys Ala Ser Gly Arg
 565 570 575
 Glu Leu Ser Tyr Val Leu Leu Ala Gly Ile Phe Leu Cys Tyr Ala Thr
 580 585 590
 Thr Phe Leu Met Ile Ala Glu Pro Asp Leu Gly Thr Cys Ser Leu Arg
 595 600 605
 Arg Ile Phe Leu Gly Leu Gly Met Ser Ile Ser Tyr Ala Ala Leu Leu
 610 615 620
 Thr Lys Thr Asn Arg Ile Tyr Arg Ile Phe Glu Gln Gly Lys Arg Ser
 625 630 635 640
 Val Ser Ala Pro Arg Phe Ile Ser Pro Ala Ser Gln Leu Ala Ile Thr
 645 650 655
 Phe Ser Leu Ile Ser Leu Gln Leu Leu Gly Ile Cys Val Trp Phe Val
 660 665 670
 Val Asp Pro Ser His Ser Val Val Asp Phe Gln Asp Gln Arg Thr Leu
 675 680 685
 Asp Pro Arg Phe Ala Arg Gly Val Leu Lys Cys Asp Ile Ser Asp Leu

690 695 700
 Ser Leu Ile Cys Leu Leu Gly Tyr Ser Met Leu Leu Met Val Thr Cys
 705 710 715 720
 Thr Val Tyr Ala Ile Lys Thr Arg Gly Val Pro Glu Thr Phe Asn Glu
 725 730 735
 Ala Lys Pro Ile Gly Phe Thr Met Tyr Thr Thr Cys Ile Val Trp Leu
 740 745 750
 Ala Phe Ile Pro Ile Phe Phe Gly Thr Ser Gln Ser Ala Asp Lys Leu
 755 760 765
 Tyr Ile Gln Thr Thr Thr Leu Thr Val Ser Val Ser Leu Ser Ala Ser
 770 775 780
 Val Ser Leu Gly Met Leu Tyr Met Pro Lys Val Tyr Ile Ile Leu Phe
 785 790 795 800
 His Pro Glu Gln Asn Val Pro Lys Arg Lys Arg Ser Leu Lys Ala Val
 805 810 815
 Val Thr Ala Ala Thr Met Ser Asn Lys Phe Thr Gln Lys Gly Asn Phe
 820 825 830
 Arg Pro Asn Gly Glu Ala Lys Ser Glu Leu Cys Glu Asn Leu Glu Ala
 835 840 845
 Pro Ala Leu Ala Thr Lys Gln Thr Tyr Val Thr Tyr Thr Asn His Ala
 850 855 860
 Ile
 865

 <210> 42
 <211> 845
 <212> PRT
 <213> Homo sapiens

 <400> 42
 Met Glu Thr Lys Gly Tyr His Ser Leu Pro Glu Gly Leu Asp Met Glu
 1 5 10 15
 Arg Arg Trp Gly Gln Val Ser Gln Ala Val Glu Arg Ser Ser Leu Gly
 20 25 30
 Pro Thr Glu Arg Thr Asp Glu Asn Asn Tyr Met Glu Ile Val Asn Val
 35 40 45
 Ser Cys Val Ser Gly Ala Ile Pro Asn Asn Ser Thr Gln Gly Ser Ser
 50 55 60
 Lys Glu Lys Gln Glu Leu Leu Pro Cys Leu Gln Gln Asp Asn Asn Arg
 65 70 75 80
 Pro Gly Ile Leu Thr Ser Asp Ile Lys Thr Glu Leu Glu Ser Lys Glu
 85 90 95
 Leu Ser Ala Thr Val Ala Glu Ser Met Gly Leu Tyr Met Asp Ser Val

100							105					110						
Arg	Asp	Ala	Asp	Tyr	Ser	Tyr	Glu	Gln	Gln	Asn	Gln	Gln	Gly	Ser	Met			
		115					120					125						
Ser	Pro	Ala	Lys	Ile	Tyr	Gln	Asn	Val	Glu	Gln	Leu	Val	Lys	Phe	Tyr			
		130				135					140							
Lys	Gly	Asn	Gly	His	Arg	Pro	Ser	Thr	Leu	Ser	Cys	Val	Asn	Thr	Pro			
145					150					155					160			
Leu	Arg	Ser	Phe	Met	Ser	Asp	Ser	Gly	Ser	Ser	Val	Asn	Gly	Gly	Val			
				165					170					175				
Met	Arg	Ala	Ile	Val	Lys	Ser	Pro	Ile	Met	Cys	His	Glu	Lys	Ser	Pro			
			180					185					190					
Ser	Val	Cys	Ser	Pro	Leu	Asn	Met	Thr	Ser	Ser	Val	Cys	Ser	Pro	Ala			
		195				200						205						
Gly	Ile	Asn	Ser	Val	Ser	Ser	Thr	Thr	Ala	Ser	Phe	Gly	Ser	Phe	Pro			
		210				215					220							
Val	His	Ser	Pro	Ile	Thr	Gln	Gly	Thr	Pro	Leu	Thr	Cys	Ser	Pro	Asn			
225					230					235					240			
Ala	Glu	Asn	Arg	Gly	Ser	Arg	Ser	His	Ser	Pro	Ala	His	Ala	Ser	Asn			
				245				250						255				
Val	Gly	Ser	Pro	Leu	Ser	Ser	Pro	Leu	Ser	Ser	Met	Lys	Ser	Ser	Ile			
			260					265					270					
Ser	Ser	Pro	Pro	Ser	His	Cys	Ser	Val	Lys	Ser	Pro	Val	Ser	Ser	Pro			
		275				280						285						
Asn	Asn	Val	Thr	Leu	Arg	Ser	Ser	Val	Ser	Ser	Pro	Ala	Asn	Ile	Asn			
		290				295						300						
Asn	Ser	Arg	Cys	Ser	Val	Ser	Ser	Pro	Ser	Asn	Thr	Asn	Asn	Arg	Ser			
305					310					315					320			
Thr	Leu	Ser	Ser	Pro	Ala	Ala	Ser	Thr	Val	Gly	Ser	Ile	Cys	Ser	Pro			
				325					330					335				
Val	Asn	Asn	Ala	Phe	Ser	Tyr	Thr	Ala	Ser	Gly	Thr	Ser	Ala	Gly	Ser			
			340					345					350					
Ser	Thr	Leu	Arg	Asp	Val	Val	Pro	Ser	Pro	Asp	Thr	Gln	Glu	Lys	Gly			
		355				360						365						
Ala	Gln	Glu	Val	Pro	Phe	Pro	Lys	Thr	Glu	Glu	Val	Glu	Ser	Ala	Ile			
						375					380							
Ser	Asn	Gly	Val	Thr	Gly	Gln	Leu	Asn	Ile	Val	Gln	Tyr	Ile	Lys	Pro			
385					390					395					400			
Glu	Pro	Asp	Gly	Ala	Phe	Ser	Ser	Ser	Cys	Leu	Gly	Gly</						

Lys His Ser Cys Ser Gly Thr Ser Phe Lys Gly Asn Pro Thr Val Asn
 435 440 445
 Pro Phe Pro Phe Met Asp Gly Ser Tyr Phe Ser Phe Met Asp Asp Lys
 450 455 460
 Asp Tyr Tyr Ser Leu Ser Gly Ile Leu Gly Pro Pro Val Pro Gly Phe
 465 470 475 480
 Asp Gly Asn Cys Glu Gly Ser Gly Phe Pro Val Gly Ile Lys Gln Glu
 485 490 495
 Pro Asp Asp Gly Ser Tyr Tyr Pro Glu Ala Ser Ile Pro Ser Ser Ala
 500 505 510
 Ile Val Gly Val Asn Ser Gly Gly Gln Ser Phe His Tyr Arg Ile Gly
 515 520 525
 Ala Gln Gly Thr Ile Ser Leu Ser Arg Ser Ala Arg Asp Gln Ser Phe
 530 535 540
 Gln His Leu Ser Ser Phe Pro Pro Val Asn Thr Leu Val Glu Ser Trp
 545 550 555 560
 Lys Ser His Gly Asp Leu Ser Ser Arg Arg Ser Asp Gly Tyr Pro Val
 565 570 575
 Leu Glu Tyr Ile Pro Glu Asn Val Ser Ser Ser Thr Leu Arg Ser Val
 580 585 590
 Ser Thr Gly Ser Ser Arg Pro Ser Lys Ile Cys Leu Val Cys Gly Asp
 595 600 605
 Glu Ala Ser Gly Cys His Tyr Gly Val Val Thr Cys Gly Ser Cys Lys
 610 615 620
 Val Phe Phe Lys Arg Ala Val Glu Gly Gln His Asn Tyr Leu Cys Ala
 625 630 635 640
 Gly Arg Asn Asp Cys Ile Ile Asp Lys Ile Arg Arg Lys Asn Cys Pro
 645 650 655
 Ala Cys Arg Leu Gln Lys Cys Leu Gln Ala Gly Met Asn Leu Gly Ala
 660 665 670
 Arg Lys Ser Lys Lys Leu Gly Lys Leu Lys Gly Ile His Glu Glu Gln
 675 680 685
 Pro Gln Gln Gln Gln Pro Pro Pro Pro Pro Pro Pro Gln Ser Pro
 690 695 700
 Glu Glu Gly Thr Thr Tyr Ile Ala Pro Ala Lys Glu Pro Ser Val Asn
 705 710 715 720
 Thr Ala Leu Val Pro Gln Leu Ser Thr Ile Ser Arg Ala Leu Thr Pro
 725 730 735
 Ser Pro Val Met Val Leu Glu Asn Ile Glu Pro Glu Ile Val Tyr Ala
 740 745 750

Gly Tyr Asp Ser Ser Lys Pro Asp Thr Ala Glu Asn Leu Leu Ser Thr
755 760 765

Leu Asn Arg Leu Ala Gly Lys Gln Met Ile Gln Val Val Lys Trp Ala
770 775 780

Lys Val Leu Pro Gly Phe Lys Asn Leu Pro Leu Glu Asp Gln Ile Thr
785 790 795 800

Leu Ile Gln Tyr Ser Trp Met Cys Leu Ser Ser Phe Ala Leu Ser Trp
805 810 815

Arg Ser Tyr Lys His Thr Asn Ser Gln Phe Leu Tyr Phe Ala Pro Asp
820 825 830

Leu Val Phe Asn Glu Leu Leu Ala Arg Val Arg Glu Gly
835 840 845

<210> 43

<211> 837

<212> PRT

<213> Homo sapiens

<400> 43

Met Glu Thr Lys Gly Tyr His Ser Leu Pro Glu Gly Leu Asp Met Glu
1 5 10 15

Arg Arg Trp Gly Gln Val Ser Cln Ala Val Glu Arg Ser Ser Leu Gly
20 25 30

Pro Thr Glu Arg Thr Asp Glu Asn Asn Tyr Met Glu Ile Val Asn Val
35 40 45

Ser Cys Val Ser Gly Ala Ile Pro Asn Asn Ser Thr Gln Gly Ser Ser
50 55 60

Lys Glu Lys Gln Glu Leu Leu Pro Cys Leu Gln Gln Asp Asn Asn Arg
65 70 75 80

Pro Gly Ile Leu Thr Ser Asp Ile Lys Thr Glu Leu Glu Ser Lys Glu
85 90 95

Leu Ser Ala Thr Val Ala Glu Ser Met Gly Leu Tyr Met Asp Ser Val
100 105 110

Arg Asp Ala Asp Tyr Ser Tyr Glu Gln Gln Asn Gln Gln Gly Ser Met
115 120 125

Ser Pro Ala Lys Ile Tyr Gln Asn Val Glu Gln Leu Val Lys Phe Tyr
130 135 140

Lys Gly Asn Gly His Arg Pro Ser Thr Leu Ser Cys Val Asn Thr Pro
145 150 155 160

Leu Arg Ser Phe Met Ser Asp Ser Gly Ser Ser Val Asn Gly Gly Val
165 170 175

Met Arg Ala Ile Val Lys Ser Pro Ile Met Cys His Glu Lys Ser Pro
180 185 190

Ser Val Cys Ser Pro Leu Asn Met Thr Ser Ser Val Cys Ser Pro Ala
 195 200 205
 Gly Ile Asn Ser Val Ser Ser Thr Thr Ala Ser Phe Gly Ser Phe Pro
 210 215 220
 Val His Ser Pro Ile Thr Gln Gly Thr Pro Leu Thr Cys Ser Pro Asn
 225 230 235 240
 Ala Glu Asn Arg Gly Ser Arg Ser His Ser Pro Ala His Ala Ser Asn
 245 250 255
 Val Gly Ser Pro Leu Ser Ser Pro Leu Ser Ser Met Lys Ser Ser Ile
 260 265 270
 Ser Ser Pro Pro Ser His Cys Ser Val Lys Ser Pro Val Ser Ser Pro
 275 280 285
 Asn Asn Val Thr Leu Arg Ser Ser Val Ser Ser Pro Ala Asn Ile Asn
 290 295 300
 Asn Ser Arg Cys Ser Val Ser Ser Pro Ser Asn Thr Asn Asn Arg Ser
 305 310 315 320
 Thr Leu Ser Ser Pro Ala Ala Ser Thr Val Gly Ser Ile Cys Ser Pro
 325 330 335
 Val Asn Asn Ala Phe Ser Tyr Thr Ala Ser Gly Thr Ser Ala Gly Ser
 340 345 350
 Ser Thr Leu Arg Asp Val Val Pro Ser Pro Asp Thr Gln Glu Lys Gly
 355 360 365
 Ala Gln Glu Val Pro Phe Pro Lys Thr Glu Glu Val Glu Ser Ala Ile
 370 375 380
 Ser Asn Gly Val Thr Gly Gln Leu Asn Ile Val Gln Tyr Ile Lys Pro
 385 390 395 400
 Glu Pro Asp Gly Ala Phe Ser Ser Ser Cys Leu Gly Gly Asn Ser Lys
 405 410 415
 Ile Asn Ser Asp Ser Ser Phe Ser Val Pro Ile Lys Gln Glu Ser Thr
 420 425 430
 Lys His Ser Cys Ser Gly Thr Ser Phe Lys Gly Asn Pro Thr Val Asn
 435 440 445
 Pro Phe Pro Phe Met Asp Gly Ser Tyr Phe Ser Phe Met Asp Asp Lys
 450 455 460
 Asp Tyr Tyr Ser Leu Ser Gly Ile Leu Gly Pro Pro Val Pro Gly Phe
 465 470 475 480
 Asp Gly Asn Cys Glu Gly Ser Gly Phe Pro Val Gly Ile Lys Gln Glu
 485 490 495
 Pro Asp Asp Gly Ser Tyr Tyr Pro Glu Ala Ser Ile Pro Ser Ser Ala
 500 505 510
 Ile Val Gly Val Asn Ser Gly Gly Gln Ser Phe His Tyr Arg Ile Gly

515	520	525
Ala Gln Gly Thr Ile Ser Leu Ser Arg Ser Ala Arg Asp Gln Ser Phe 530 535 540		
Gln His Leu Ser Ser Phe Pro Pro Val Asn Thr Leu Val Glu Ser Trp 545 550 555 560		
Lys Ser His Gly Asp Leu Ser Ser Arg Arg Ser Asp Gly Tyr Pro Val 565 570 575		
Leu Glu Tyr Ile Pro Glu Asn Val Ser Ser Ser Thr Leu Arg Ser Val 580 585 590		
Ser Thr Gly Ser Ser Arg Pro Ser Lys Ile Cys Leu Val Cys Gly Asp 595 600 605		
Glu Ala Ser Gly Cys His Tyr Gly Val Val Thr Cys Gly Ser Cys Lys 610 615 620		
Val Phe Phe Lys Arg Ala Val Glu Gly Gln His Asn Tyr Leu Cys Ala 625 630 635 640		
Gly Arg Asn Asp Cys Ile Ile Asp Lys Ile Arg Arg Lys Asn Cys Pro 645 650 655		
Ala Cys Arg Leu Gln Lys Cys Leu Gln Ala Gly Met Asn Leu Gly Ala 660 665 670		
Arg Lys Ser Lys Lys Leu Gly Lys Leu Lys Gly Ile His Glu Glu Gln 675 680 685		
Pro Gln Gln Gln Gln Pro Pro Pro Pro Pro Pro Pro Gln Ser Pro 690 695 700		
Glu Glu Gly Thr Thr Tyr Ile Ala Pro Ala Lys Glu Pro Ser Val Asn 705 710 715 720		
Thr Ala Leu Val Pro Gln Leu Ser Thr Ile Ser Arg Ala Leu Thr Pro 725 730 735		
Ser Pro Val Met Val Leu Glu Asn Ile Glu Pro Glu Ile Val Tyr Ala 740 745 750		
Gly Tyr Asp Ser Ser Lys Pro Asp Thr Ala Glu Asn Leu Leu Ser Thr 755 760 765		
Leu Asn Arg Leu Ala Gly Lys Gln Met Ile Gln Val Val Lys Trp Ala 770 775 780		
Lys Val Leu Pro Gly Phe Lys Asn Leu Pro Leu Glu Asp Gln Ile Thr 785 790 795 800		
Leu Ile Gln Tyr Ser Trp Met Cys Leu Ser Ser Phe Ala Leu Ser Trp 805 810 815		
Arg Ser Tyr Lys His Thr Asn Ser Gln Phe Leu Tyr Phe Ala Pro Asp 820 825 830		
Leu Val Phe Asn Glu 835		

<210> 44
 <211> 640
 <212> PRT
 <213> Homo sapiens

<400> 44

Met Gly Arg Leu Gln Leu Val Val Leu Gly Leu Thr Cys Cys Trp Ala
 1 5 10 15

Val Ala Ser Ala Ala Lys Leu Gly Ala Val Tyr Thr Glu Gly Gly Phe
 20 25 30

Val Glu Gly Val Asn Lys Lys Leu Gly Leu Leu Gly Asp Ser Val Asp
 35 40 45

Ile Phe Lys Gly Ile Pro Phe Ala Ala Pro Thr Lys Ala Leu Glu Asn
 50 55 60

Pro Gln Pro His Pro Gly Trp Gln Gly Thr Leu Lys Ala Lys Asn Phe
 65 70 75 80

Lys Lys Arg Cys Leu Gln Ala Thr Ile Thr Gln Asp Ser Thr Tyr Gly
 85 90 95

Asp Glu Asp Cys Leu Tyr Leu Asn Ile Trp Val Pro Gln Gly Arg Lys
 100 105 110

Gln Val Ser Arg Asp Leu Pro Val Met Ile Trp Ile Tyr Gly Gly Ala
 115 120 125

Phe Leu Met Gly Ser Gly His Gly Ala Asn Phe Leu Asn Asn Tyr Leu
 130 135 140

Tyr Asp Gly Glu Glu Ile Ala Thr Arg Gly Asn Val Ile Val Val Thr
 145 150 155 160

Phe Asn Tyr Arg Val Gly Pro Leu Gly Phe Leu Ser Thr Gly Asp Ala
 165 170 175

Asn Leu Pro Gly Asn Tyr Gly Leu Arg Asp Gln His Met Ala Ile Ala
 180 185 190

Trp Val Lys Arg Asn Ile Ala Ala Phe Gly Gly Asp Pro Asn Asn Ile
 195 200 205

Thr Leu Phe Gly Glu Ser Ala Gly Gly Ala Ser Val Ser Leu Gln Thr
 210 215 220

Leu Ser Pro Tyr Asn Lys Gly Leu Ile Arg Arg Ala Ile Ser Gln Ser
 225 230 235 240

Gly Val Ala Leu Ser Pro Trp Val Ile Gln Lys Asn Pro Leu Phe Trp
 245 250 255

Ala Lys Lys Val Ala Glu Lys Val Gly Cys Pro Val Gly Asp Ala Ala
 260 265 270

Arg Met Ala Gln Cys Leu Lys Val Thr Asp Pro Arg Ala Leu Thr Leu
 275 280 285

Ala Tyr Lys Val Pro Leu Ala Gly Leu Glu Tyr Pro Met Leu His Tyr
 290 295 300
 Val Gly Phe Val Pro Val Ile Asp Gly Asp Phe Ile Pro Ala Asp Pro
 305 310 315 320
 Ile Asn Leu Tyr Ala Asn Ala Ala Asp Ile Asp Tyr Ile Ala Gly Thr
 325 330 335
 Asn Asn Met Asp Gly His Ile Phe Ala Ser Ile Asp Met Pro Ala Ile
 340 345 350
 Asn Lys Gly Asn Lys Lys Val Thr Glu Glu Asp Phe Tyr Lys Leu Val
 355 360 365
 Ser Glu Phe Thr Ile Thr Lys Gly Leu Arg Gly Ala Lys Thr Thr Phe
 370 375 380
 Asp Val Tyr Thr Glu Ser Trp Ala Gln Asp Pro Ser Gln Glu Asn Lys
 385 390 395 400
 Lys Lys Thr Val Val Asp Phe Glu Thr Asp Val Leu Phe Leu Val Pro
 405 410 415
 Thr Glu Ile Ala Leu Ala Gln His Arg Ala Asn Ala Lys Ser Ala Lys
 420 425 430
 Thr Tyr Ala Tyr Leu Phe Ser His Pro Ser Arg Met Pro Val Tyr Pro
 435 440 445
 Lys Trp Val Gly Ala Asp His Ala Asp Asp Ile Gln Tyr Val Phe Gly
 450 455 460
 Lys Pro Phe Ala Thr Pro Thr Gly Tyr Arg Pro Gln Asp Arg Thr Val
 465 470 475 480
 Ser Lys Ala Met Ile Ala Tyr Trp Thr Asn Phe Ala Lys Thr Gly Asp
 485 490 495
 Pro Asn Met Gly Asp Ser Ala Val Pro Thr His Trp Glu Pro Tyr Thr
 500 505 510
 Thr Glu Asn Ser Gly Tyr Leu Glu Ile Thr Lys Lys Met Gly Ser Ser
 515 520 525
 Ser Met Lys Arg Ser Leu Arg Thr Asn Phe Leu Arg Tyr Trp Thr Leu
 530 535 540
 Thr Tyr Leu Ala Leu Pro Thr Val Thr Asp Gln Glu Ala Thr Pro Val
 545 550 555 560
 Pro Pro Thr Gly Asp Ser Glu Ala Thr Pro Val Pro Pro Thr Gly Asp
 565 570 575
 Ser Glu Thr Ala Pro Val Pro Pro Thr Gly Asp Ser Gly Ala Pro Pro
 580 585 590
 Val Pro Pro Thr Gly Asp Ser Gly Ala Pro Pro Val Pro Pro Thr Gly
 595 600 605

Asp Ser Gly Ala Pro Pro Val Pro Pro Thr Gly Asp Ser Gly Ala Pro
610 615 620

Pro Val Pro Pro Thr Gly Asp Ser Gly Ala Pro Pro Val Pro Pro Pro
625 630 635 640

<210> 45
<211> 201
<212> PRT
<213> Homo sapiens

<400> 45
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1 5 10 15
Asp Ser Lys Gly Ser Asn Glu Leu His Gln Val Pro Ser Asn Cys Asp
20 25 30
Cys Leu Asn Gly Gly Thr Cys Val Ser Asn Lys Tyr Phe Ser Asn Ile
35 40 45
His Trp Cys Asn Cys Pro Lys Lys Phe Gly Gly Gln His Cys Glu Ile
50 55 60
Asp Lys Ser Lys Thr Cys Tyr Glu Gly Asn Gly His Phe Tyr Arg Gly
65 70 75 80
Lys Ala Ser Thr Asp Thr Met Gly Arg Pro Cys Leu Pro Trp Asn Ser
85 90 95
Ala Thr Val Leu Gln Gln Thr Tyr His Ala His Arg Ser Asp Ala Leu
100 105 110
Gln Leu Gly Leu Gly Lys His Asn Tyr Cys Arg Glu Val Gly Ala Gln
115 120 125
Gly Pro Lys Ala Leu Pro Thr Val Pro Arg Asn Leu Val Thr Ile Pro
130 135 140
Phe Ser Gln Arg Ala Gly His Ser Thr Arg Glu Val Gln Pro Leu Val
145 150 155 160
Glu Ser Ser Leu Arg Gly Gly Gly Arg Glu Gly Pro Leu Gly Trp Asn
165 170 175
Asp Ile Pro Tyr Leu Ser Val Leu Pro Gly Thr Gln Thr Thr Gly Gly
180 185 190
Asp Pro Gly Ala Met Cys Arg Trp Ala
195 200

<210> 46
<211> 74
<212> PRT
<213> Homo sapiens

<400> 46

Met Lys Thr Tyr Arg Ala Lys Phe Cys Gly Val Cys Thr Asp Gly Arg
 1 5 10 15

Cys Cys Thr Pro His Arg Thr Thr Thr Leu Pro Val Glu Phe Lys Cys
 20 25 30

Pro Asp Gly Glu Val Met Lys Lys Asn Met Met Phe Ile Lys Thr Cys
 35 40 45

Ala Cys His Tyr Asn Cys Pro Gly Asp Asn Asp Ile Phe Glu Ser Leu
 50 55 60

Tyr Tyr Arg Lys Met Tyr Gly Asp Met Ala
 65 70

<210> 47

<211> 166

<212> PRT

<213> Homo sapiens

<400> 47

Met Thr Ala Ala Ser Met Gly Pro Val Arg Val Ala Phe Val Val Leu
 1 5 10 15

Leu Ala Leu Cys Ser Arg Pro Ala Val Gly Gln Asn Cys Ser Gly Pro
 20 25 30

Cys Arg Cys Pro Asp Glu Pro Ala Pro Arg Cys Pro Ala Gly Val Ser
 35 40 45

Leu Val Leu Asp Gly Cys Gly Cys Cys Arg Val Cys Ala Lys Gln Leu
 50 55 60

Gly Glu Leu Cys Thr Glu Arg Asp Pro Cys Asp Pro His Lys Gly Leu
 65 70 75 80

Phe Cys Asp Phe Gly Ser Pro Ala Asn Arg Lys Ile Gly Val Cys Thr
 85 90 95

Ala Lys Asp Gly Ala Pro Cys Ile Phe Gly Gly Thr Val Tyr Arg Ser
 100 105 110

Gly Glu Ser Phe Gln Ser Ser Cys Lys Tyr Gln Cys Thr Cys Leu Asp
 115 120 125

Gly Ala Val Gly Cys Met Pro Leu Cys Ser Met Asp Val Arg Leu Pro
 130 135 140

Ser Pro Asp Cys Pro Phe Pro Ser Leu Pro Thr Gly Arg His Val Trp
 145 150 155 160

Pro Arg Pro Asn Tyr Asp
 165

<210> 48

<211> 140

<212> PRT

<213> Homo sapiens

<400> 48

Met Glu Asn Ser Leu Arg Cys Val Trp Val Pro Lys Leu Ala Phe Val
1 5 10 15

Leu Phe Gly Ala Ser Leu Leu Ser Ala His Leu Gln Val Thr Gly Phe
20 25 30

Gln Ile Lys Ala Phe Thr Ala Leu Arg Phe Leu Ser Glu Pro Ser Asp
35 40 45

Ala Val Thr Met Arg Gly Gly Asn Val Leu Leu Asp Cys Ser Ala Glu
50 55 60

Ser Asp Arg Gly Val Pro Val Ile Lys Trp Lys Lys Asp Cys Ile His
65 70 75 80

Leu Ala Leu Gly Met Asp Glu Arg Lys Gln Gln Leu Ser Asn Gly Ser
85 90 95

Leu Leu Ile Gln Asn Ile Leu His Ser Arg His His Lys Pro Asp Glu
100 105 110

Gly Leu Tyr Gln Cys Glu Ala Ser Leu Gly Asp Ser Gly Ser Ile Ile
115 120 125

Ser Arg Thr Ala Lys Val Ala Val Ala Gly Pro Thr
130 135 140

<210> 49

<211> 147

<212> PRT

<213> Homo sapiens

<400> 49

Met Ala Pro Phe Glu Pro Leu Ala Ser Gly Ile Leu Leu Leu Trp
1 5 10 15

Leu Ile Ala Pro Ser Arg Ala Cys Thr Cys Val Pro Pro His Pro Gln
20 25 30

Thr Ala Phe Cys Asn Ser Asp Leu Val Ile Arg Ala Lys Phe Val Gly
35 40 45

Thr Pro Gln Val Asn Gln Thr Thr Leu Tyr Gln Arg Tyr Glu Ile Lys
50 55 60

Met Thr Lys Met Tyr Lys Gly Phe Gln Ala Leu Xaa Asp Ala Ala Asp
65 70 75 80

Ile Arg Phe Val Tyr Thr Pro Ala Met Glu Ser Val Cys Gly Tyr Phe
85 90 95

His Arg Ser His Asn Arg Ser Arg Gly Val Ser His Cys Trp Lys Thr
100 105 110

Ala Gly Trp Thr Leu Ala His His Tyr Leu Gln Phe Arg Gly Ser Leu
115 120 125

Glu Gln Pro Glu Leu Ser Ser Ala Pro Gly Leu His Gln Asp Leu His

130 135 140

Cys Trp Leu
145

<210> 50
<211> 82
<212> PRT
<213> Homo sapiens

<400> 50
Met Ser Val Lys Glu Thr Leu Pro Leu Ile His Gln Gln Met Tyr Lys
1 5 10 15

Gly Phe Gln Ala Leu Gly Asp Ala Ala Asp Ile Arg Phe Val Tyr Thr
20 25 30

Pro Ala Met Glu Ser Val Cys Gly Tyr Phe His Arg Ser His Asn Arg
35 40 45

Ser Glu Glu Phe Leu Ile Ala Gly Glu Ala Pro Ser Pro Arg Pro Val
50 55 60

Pro His Gln Pro Val Pro Gly Ala Arg Pro Ser Asn His Glu Gly Ala
65 70 75 80

Arg Leu

<210> 51
<211> 115
<212> PRT
<213> Homo sapiens

<400> 51
Met Ala Pro Phe Glu Pro Leu Ala Ser Gly Ile Leu Leu Leu Trp
1 5 10 15

Leu Ile Ala Pro Ser Arg Ala Cys Thr Cys Val Pro Pro His Pro Gln
20 25 30

Thr Ala Phe Cys Asn Ser Asp Leu Val Ile Arg Ala Lys Phe Val Gly
35 40 45

Thr Pro Glu Val Asn Gln Thr Thr Leu Tyr Gln Arg Tyr Glu Ile Lys
50 55 60

Met Thr Lys Met Tyr Lys Gly Phe Gln Ala Leu Gly Asp Ala Ala Asp
65 70 75 80

Ile Arg Phe Val Tyr Thr Pro Ala Met Glu Ser Val Cys Gly Tyr Phe
85 90 95

His Arg Met Asp Ser Cys Thr Ser Leu Pro Ala Val Ser Trp Leu Pro
100 105 110

Gly Thr Ala
115

<210> 52
 <211> 143
 <212> PRT
 <213> Homo sapiens

<400> 52
 Met Thr Lys Met Tyr Lys Gly Phe Gln Ala Leu Gly Asp Ala Ala Asp
 1 5 10 15
 Ile Arg Phe Val Tyr Thr Pro Ala Met Glu Ser Val Cys Gly Tyr Phe
 20 25 30
 His Arg Ser His Asn Arg Ser Glu Glu Phe Leu Ile Ala Gly Lys Leu
 35 40 45
 Gln Asp Gly Leu Leu His Ile Thr Thr Cys Ser Phe Val Ala Pro Trp
 50 55 60
 Asn Ser Leu Ser Leu Ala Gln Arg Arg Gly Phe Thr Lys Thr Tyr Thr
 65 70 75 80
 Val Gly Cys Glu Glu Cys Thr Val Phe Pro Cys Leu Ser Ile Pro Cys
 85 90 95
 Lys Leu Gln Ser Gly Thr His Cys Leu Trp Thr Asp Gln Leu Leu Gln
 100 105 110
 Gly Ser Glu Lys Gly Phe Gln Ser Arg His Leu Ala Cys Leu Pro Arg
 115 120 125
 Glu Pro Gly Leu Cys Thr Trp Gln Ser Leu Arg Ser Gln Ile Ala
 130 135 140

1 MRENMARGPCNAPRWVSLMVLVAIGTAVTAAVNPVVVRISQKGLDYASQ 50
|||||
1 MRENMARGPCNAPRWVSLMVLVAIGTAVTAAVNPVVVRISQKGLDYASQ 50

51 QGTAALQKELKRIKIPDYSDFKIKHLGKGHYSFYSMDIRFQLPSSQIS 100
|||||
51 QGTAALQKELKRIKIPDYSDFKIKHLGKGHYSFYSMDIRFQLPSSQIS 100

101 MVPNVGLKFESISNANIKISGKWKAKRFLKMSGNFDLSIEGMSISADLKL 150
|||||
101 MVPNVGLKFESISNANIKISGKWKAKRFLKMSGNFDLSIEGMSISADLKL 150

151 GSNPTSGKPTITCSCSSHINSVHVHISKSKVGWLIQLFHKKIESALRNK 200
|||||
151 GSNPTSGKPTITCSCSSHINSVHVHISKSKVGWLIQLFHKKIESALRNK 200

201 MNSQVCEKVTNSVSSKLQPYFQTLFVMTKIDSVAGINYLVAAPPATTAET 250
|||||
201 MNSQVCEKVTNSVSSKLQPYFQTLFVMTKIDSVAGINYLVAAPPATTAET 250

Fig. 1

251 LDVQMKGEFYSENHHNPPPFAPPVMEFFAAHDMVYLGLSDYFFNTAGLV 300
|||||
251 LDVQMKGEFYSENHHNPPPFAPPVMEFFAAHDMVYLGLSDYFFNTAGLV 300

301 YQEAGVLKMTLRDDMIPKESKERLTTKFFGTFLPEVAKKFPNMKIQIHVS 350
|||||
301 YQEAGVLKMTLRDDMIPKESKERLTTKFFGTFLPEVAKKFPNMKIQIHVS 350

351 ASTPPHLSVQPTGLTFYPADVQAFVLPNSSLASLFLIGM 391
|||||
351 ASTPPHLSVQPTGLTFYPADVQAFVLPNSSLASLFLIGM 391

Fig. 1 (Cont.)

1 MGRQLVLVLGLTCCWAVASAAKLGAVYTEGGFVEGVNKKLGLLGDSDVDF 50
|||||
1 MGRQLVLVLGLTCCWAVASAAKLGAVYTEGGFVEGVNKKLGLLGDSDVDF 50
51 KGIPFAAPTCALENPQHPGWQGTAKAKNEKKRCLQATITQDSTYGDED 100
|||||
51 KGIPFAAPTCALENPQHPGWQGTAKAKNEKKRCLQATITQDSTYGDED 100
101 LYLNIWVVPQGRKQVSRDLPVMIWIYGGAFMGSGHGANELNNLYLDGEEI 150
|||||
101 LYLNIWVVPQGRKQVSRDLPVMIWIYGGAFMGSGHGANELNNLYLDGEEI 150
151 ATRGNVIVVTENYRVGPLGFLSTGDANLPGNYGLRDQHMAIAWVKRNIAA 200
|||||
151 ATRGNVIVVTENYRVGPLGFLSTGDANLPGNYGLRDQHMAIAWVKRNIAA 200
201 FGGDPNNITLFGEAGGASVSLQTLSPYNKGLIRRAISQSGVALSPWVIQ 250
|||||
201 FGGDPNNITLFGEAGGASVSLQTLSPYNKGLIRRAISQSGVALSPWVIQ 250

Fig. 2

251 KNPLFWAKKVAEKVGCPCVGDAARMAQCLKVTDPPRALTLAYKVPLAGLEYP 300
|||||
251 KNPLFWAKKVAEKVGCPCVGDAARMAQCLKVTDPPRALTLAYKVPLAGLEYP 300
301 MLHYVGFVPVIDGDFIPADPINLYANAADIDYIAGTNMMDGHIFASIDMP 350
|||||
301 MLHYVGFVPVIDGDFIPADPINLYANAADIDYIAGTNMMDGHIFASIDMP 350
351 AINKGNKKVTEEDFYKLVSEFTITKGLRGAKTTFDVYTESWAQDPSQENK 400
|||||
351 AINKGNKKVTEEDFYKLVSEFTITKGLRGAKTTFDVYTESWAQDPSQENK 400
401 KKTVDVDETDVLFVLPTEIALAQHRANAKSAKTYAYLFESHPSRMPVYPKW 450
|||||
401 KKTVDVDETDVLFVLPTEIALAQHRANAKSAKTYAYLFESHPSRMPVYPKW 450
451 VGADHADDIQYVFGKPFATPTGYRPPQDRFTVSKAMIAYWTFNFAKTGDPNMG 500
|||||
451 VGADHADDIQYVFGKPFATPTGYRPPQDRFTVSKAMIAYWTFNFAKTGDPNMG 500

Fig. 2 (Cont.)

501 DSAVPTHWEPTYTENS GYLEITKKMGSSSMKRSLRTNFLRYWTLTYLALP 550
|||||
501 DSAVPTHWEPTYTENS GYLEITKKMGSSSMKRSLRTNFLRYWTLTYLALP 550
551 TVTDQEATPVPPPTGDS EATPVPPPTGDS EATPVPPPTGDS GAPPVPPPTGDS G 600
|||||
551 TVTDQEATPVPPPTGDS EATPVPPPTGDS EATPVPPPTGDS GAPPVPPPTGDS G 600
601 APPVPPPTGDS GAPPVPPPTGDS SEA 623
|||||
601 APPVPPPTGDS GAPPVPPPTGDS GA 623

Fig. 2 (Cont.)

```

1  LLLGFLVSLSTLSIPPWEAPKEHKYKAEHTVVLTVTGEPCHFPEQY 50
   |||||
4  LLLGFLVSLSTLSIPPWEAPKEHKYKAEHTVVLTVTGEPCHFPEQY 53
   |||||

51 HRQLYHKCTHKGRPGPQPWCATTPNFDQDQRWGYCLEPKKVKDHCSP 100
   |||||
54 HRQLYHKCTHKGRPGPQPWCATTPNFDQDQRWGYCLEPKKVKDHCSP 103
   |||||

101 CQKGGTCVNMPSGPHCLCPQHLTGNHCQKEKCFEPQLLRFHKNWIYRT 150
   |||||
104 CQKGGTCVNMPSGPHCLCPQHLTGNHCQKEKCFEPQLLRFHKNWIYRT 153
   |||||

151 EQAAVARCQCKGPDHAHCQRLASQACRTNPCLHGRCLEVEGHRCLCHCPVG 200
   |||||
154 EQAAVARCQCKGPDHAHCQRLASQACRTNPCLHGRCLEVEGHRCLCHCPVG 203
   |||||

201 YTGPFCDVDTKASCYDGRGLSYRGLARTTLSGAPCQPWASEATYRNVTA 250
   |||||
204 YTGPFCDVDTKASCYDGRGLSYRGLARTTLSGAPCQPWASEATYRNVTA 253

```

Fig. 3

251 QARNWGLGHAFCRNPNDNDIRPWCFLNRDRLSWEYCDLAQCQTPTQAAP 300
|||||
254 QARNWGLGHAFCRNPNDNDIRPWCFLNRDRLSWEYCDLAQCQTPTQAAP 303
|||||
301 PTPVSPRLHVPLMPAQAPPKPQPTTRTPPQSQTTPGALPAKREQPPSLTR 350
|||||
304 PTPVSPRLHVPLMPAQAPPKPQPTTRTPPQSQTTPGALPAKREQPPSLTR 353
|||||
351 NGPLSCGQRLRKSLSSMTRVVGGLVALRGAHPIYAALYWGHSEFCAGSLIA 400
|||||
354 NGPLSCGQRLRKSLSSMTRVVGGLVALRGAHPIYAALYWGHSEFCAGSLIA 403
|||||
401 PCWVLTAAHCLQDRPAPEDLTVVLGQERRNHSCEPCQTLAVRSYRLHEAF 450
|||||
404 PCWVLTAAHCLQDRPAPEDLTVVLGQERRNHSCEPCQTLAVRSYRLHEAF 453
|||||
451 SPVSYQHDLALLRLQEDADGSCALLSPYVQPVCLPSGAARPSETTLCQVA 500
|||||
454 SPVSYQHDLALLRLQEDADGSCALLSPYVQPVCLPSGAARPSETTLCQVA 503
|||||

Fig. 3 (Cont.)

501 GWGHQFECAEEYASFLQEAQVPFLSLERCSAPDVHGGSSILPGMLCAGFLE 550
|||||
504 GWGHQFECAEEYASFLQEAQVPFLSLERCSAPDVHGGSSILPGMLCAGFLE 553
551 GGTDACAGELLAGWRPSPRPSSAXSQVHSADCVFPTQGDGGPLVCEQAA 600
|||||
554 GGTDAC.....QGDGGPLVCEQAA 574
601 ERRLLTQGIISWGGCGDRNKPVGVTDVAYYLAWIREHTVS 641
|||||
575 ERRLLTQGIISWGGCGDRNKPVGVTDVAYYLAWIREHTVS 615

Fig. 3 (Cont.)

```

1  MAPFEPLASGILLIWLIIAPSRACCTCPPHPQTAFCNDDLVIKAFVGTGTP 50
   |||||
1  MAPFEPLASGILLIWLIIAPSRACCTCPPHPQTAFCNDDLVIKAFVGTGTP 50
   |||||

51  EVNQTTLYQRYEIKMTKMYKGFQALGDAADIRFVYTPAMESVCGYFHRSH 100
   |||||
51  EVNQTTLYQRYEIKMTKMYKGFQALGDAADIRFVYTPAMESVCGYFHRSH 100
   |||||

101 NRSEEFLLIAGKLLQDGLLHITTCSEFVAPWNSLSLAQRRGFTKTYTVGCCEC 150
   |||||
101 NRSEEFLLIAGKLLQDGLLHITTCSEFVAPWNSLSLAQRRGFTKTYTVGCCEC 150
   |||||

151 TVFPC 155
   |||||
151 TVFPC 155

```

Fig. 4

[illegible]

Fig. 5

```

1 MAPFEPLASGILLLLWLIAPSRACTCVPPHPQTAFNCSDLVIRAKFVGTP 50
  |||||
1 MAPFEPLASGILLLLWLIAPSRACTCVPPHPQTAFNCSDLVIRAKFVGTP 50

51 EVNQTTIYQRYEIKMTKMYKGFQALGDAADIRFVYTPAMESVCGYFHRSH 100
  |||||
51 EVNQTTIYQRYEIKMTKMYKGFQALGDAADIRFVYTPAMESVCGYFHRSH 100

101 NRSEEFLLLGKLQDGFIAHSLTCSFCWVPWENSLSLAQRRGFTKTYTVG 150
  ||||| : |||||
101 NRSEEFLLI.AGKLQDGLL.HITCSFV.APW.NSLSLAQRRGFTKTYTVG 146

151 CEECTVFPCLSI PCKLQSGTHCLWTDQLLQSEKGFQSRHLLACLPREPGL 200
  |||||
147 CEECTVFPCLSI PCKLQSGTHCLWTDQLLQSEKGFQSRHLLACLPREPGL 196

201 CTWQSLRSQIA 211
  |||||
197 CTWQSLRSQIA 207

```

Fig. 6

```

1  MAPFEPLASGILLLLWLIAPSRACCTCVPPHPQTAFCNSDLVIRAKFVGTP 50
   |||||
1  MAPFEPLASGILLLLWLIAPSRACCTCVPPHPQTAFCNSDLVIRAKFVGTP 50
   |||||
51 EVNQTTLYQRYEIKMTKMYKGFQALGDAADIRFVYTPAMESVCGYFHRSH 100
   |||||
51 EVNQTTLYQRYEIKMTKMYKGFQALGDAADIRFVYTPAMESVCGYFHRSH 100
   |||||
101 NRSEEEFLI..... 108
     |||||
101 NRSEEEFLIAGKIQDGLLHITTCSEFVAPWNLSLAQRRGFTTKYTVGCCEC 150
   .
109 .....LSIPCKLQSGTHCLWTDQLLQGSEKGFQSRHLACLPREPGLCTWQ 153
     |||||
151 TVFPCLSIIPCKLQSGTHCLWTDQLLQGSEKGFQSRHLACLPREPGLCTWQ 200
   .
   154 SLRSQIA 160
     |||||
   201 SLRSQIA 207

```

Fig. 7

```

1  MAPFEPLASGILLLLWLIAPSRACCTCVPPHPQTAFCNSDLVIRAKFVGTP 50
   |||||
1  MAPFEPLASGILLLLWLIAPSRACCTCVPPHPQTAFCNSDLVIRAKFVGTP 50
   |||||

51  EVNQTTLYQRYEIKMTKMYKGFQALGDAADIRFVYTPAMESVCGYFHR.. 98
   |||||
51  EVNQTTLYQRYEIKMTKMYKGFQALGDAADIRFVYTPAMESVCGYFHRSH 100
   |||||

99  .....AGKLQDGLLHITTCSFVAPWNSLSLAQRRGFTKTYTVGCEE 140
   |||||
101  NRSEEFLLAGKLQDGLLHITTCSFVAPWNSLSLAQRRGFTKTYTVGCEE 150
   |||||

141  TVFPCLSI PCKLQSGTHCLWTDQLLQSGEKGFQSRHLACLPREPGLCTWQ 190
   |||||
151  TVFPCLSI PCKLQSGTHCLWTDQLLQSGEKGFQSRHLACLPREPGLCTWQ 200
   |||||

191  SLRSQIA 197
    |||||
201  SLRSQIA 207

```

Fig. 8

Fig. 6

251 ENQPWFAAIYRRHRGGSVTYVCGGSLISPCWVISATHCFIDYPKKEDYIV 300
 |||||
 188 ENQPWFAAIYRRHRGGSVTYVCGGSLISPCWVISATHCFIDYPKKEDYIV 237
 |||||
 301 YLGRSRLNSNTQGENKFEVENLILHKDYSADTLAHNDIALLKIRSEGR 350
 |||||
 238 YLGRSRLNSNTQGENKFEVENLILHKDYSADTLAHNDIALLKIRSEGR 287
 |||||
 351 CAQPSRTIQTICLPSMYNDPQFGTSCEITGFGKENSTDYLYPEQLKMTVV 400
 |||||
 288 CAQPSRTIQTICLPSMYNDPQFGTSCEITGFGKENSTDYLYPEQLKMTVV 337
 |||||
 401 KLISHRECQQPHYYGSEVTTKMLCAADPQWKTDSCQGDGSGGPLVCSLQGR 450
 |||||
 338 KLISHRECQQPHYYGSEVTTKMLCAADPQWKTDSCQGDGSGGPLVCSLQCR 387
 |||||
 451 MTLTGIVSWGRCALKDKPGVYTRVSHFLPWIRSHTKKEENGLAL 494
 |||||
 388 MTLTGIVSWGRCALKDKPGVYTRVSHFLPWIRSHTKKEENGLVL 431

Fig. 9 (Cont.)

```

1  MQMSPALTCVLGLALVFEGESAVHHPPSYVAHLASDFGVRVFQQAQAS 50
   |||||
1  MQMSPALTCVLGLALVFEGESAVHHPPSYVAHLASDFGVRVFQQAQAS 50

51  KDRNVVFSFYGVASVLAMLQLTTGGETQQQIQAAAMGFKIDDKGMAPALRH 100
   |||||
51  KDRNVVFSFYGVASVLAMLQLTTGGETQQQIQAAAMGFKIDDKGMAPALRH 100

101 LYKELMGPWNKDEISTDAIFVQORDLKL VQGFMPPHFRLFRSTVKQVDFS 150
   |||||
101 LYKELMGPWNKDEISTDAIFVQORDLKL VQGFMPPHFRLFRSTVKQVDFS 150

151 EVERARFIINDWVKHTKGMISNLLGKGAVDQLTRLVLVNALYFNGQWKT 200
   |||||
151 EVERARFIINDWVKHTKGMISNLLGKGAVDQLTRLVLVNALYFNGQWKT 200

201 PFPDSSTHRRRLFHKSDGSTVSVPMMAQTNKFNYTEFTPDGHYYDILELP 250
   |||||
201 PFPDSSTHRRRLFHKSDGSTVSVPMMAQTNKFNYTEFTPDGHYYDILELP 250

251 YHGDTLMSMFIAADL...VPTEAL 270
   |||||
251 YHGDTLMSMFIAAPYEKEVPLSAL 273

```

Fig. 10

Fig. 11

251 YHGDTLSMFIAAPYEKEVPLSALTNIILSAQLISHWKGNMTRLPRLLVLPK 300
|||||
251 YHGDTLSMFIAAPYEKEVPLSALTNIILSAQLISHWKGNMTRLPRLLVLPK 300
.
301 FSLETEVDLRKPLENLGMDTDFRQFQADFTSLSDQEEPLHVAQALQVKIE 350
|||||
301 FSLETEVDLRKPLENLGMDTDFRQFQADFTSLSDQEEPLHVAQALQVKIE 350
.
351 VNESGTVASSSTAVIVSARMAPEEIIMDRPFLFVV 385
|||||
351 VNESGTVASSSTAVIVSARMAPEEIIMDRPFLFVV 385

Fig. 11 (Cont.)

[illegible][illegible]

169 VQTTWSACSKTCGMGISTRVTNDNASCRLEKQSRLCMVVRPCEADLEENI 218
|||||
201 VQTTWSACSKTCGMGISTRVTNDNASCRLEKQSRLCMVVRPCEADLEENI 250
|||||
219 KKGKKCIRTPKISKPIKFELSGCTSMKTYRAKFCGVCTDGRCCCTPHRTTT 268
|||||
251 KKGKKCIRTPKISKPIKFELSGCTSMKTYRAKFCGVCTDGRCCCTPHRTTT 300
|||||
269 LPVEFKCPDGEVMMKKNNMFIKTCACHYNCPGDNDIFESLYYRKMYGDMA 317
|||||
301 LPVEFKCPDGEVMMKKNNMFIKTCACHYNCPGDNDIFESLYYRKMYGDMA 349
|||||

Fig. 12 (Cont.)

Fig. 13

253 AIEGKDAVLECCVSGYPPPSFTWLRGEEVIQLRSKKYSLGGSNLLISNV 302
|||||
251 AIEGKDAVLECCVSGYPPPSFTWLRGEEVIQLRSKKYSLGGSNLLISNV 300
303 TDDDSGMYTCVVVTKYKNENISASAELTVLV 331
|||||
301 TDDDSGMYTCVVVTKYKNENISASAELTVLV 329

Fig. 13 (Cont.)

Fig. 14

303 TDDDSGMYTCVVYKNNENISASAEITVLVPPWFLNHPNLSLYAYESMDIEF 352
|||||
301 TDDDSGMYTCVVYKNNENISASAEITVLVPPWFLNHPNLSLYAYESMDIEF 350
|||||
353 ECTVSGKPVPTVNMKNGDVVIPSDYFQIVGGSNLRILGVKSDGEFYQC 402
|||||
351 ECTVSGKPVPTVNMKNGDVVIPSDYFQIVGGSNLRILGVKSDGEFYQC 400
|||||
403 VAENEAGNAQTSACLIVPKPAIPSSSVLPAPRDVVPVLVSSRFVRLSWR 452
|||||
401 VAENEAGNAQTSACLIVPKPAIPSSSVLPAPRDVVPVLVSSRFVRLSWR 450
|||||
453 PPAEAKGNIQTFTVFFSREGDNRRERALNTQPGSLQLTVGNLKPPEAMYTF 502
|||||
451 PPAEAKGNIQTFTVFFSREGDNRRERALNTQPGSLQLTVGNLKPPEAMYTF 500
|||||
503 RVVAYNEWGPGCESSQPIKVATQPELQVPGPVENLQAVSTSPSTSIITWEP 552
|||||
501 RVVAYNEWGPGCESSQPIKVATQPELQVPGPVENLQAVSTSPSTSIITWEP 550
|||||
553 PAYANGPVQGYRLFCFTEVSTGKEQNIQVDFGLSYKLEGLKKFTEYSLRFLA 602
|||||
551 PAYANGPVQGYRLFCFTEVSTGKEQNIQVDFGLSYKLEGLKKFTEYSLRFLA 600
|||||

Fig. 14 (Cont.)

Fig. 14 (Cont.)

Fig. 14 (Cont.)

Fig. 14 (Cont.)

Fig. 14 (Cont.)

1 MPGKRGLGWWWARLLPLCLLLSLYGPWMPSSLGPKPKGHPHMNSIRIDGIT 50
|||||
1 MPGKRGLGWWWARLLPLCLLLSLYGPWMPSSLGPKPKGHPHMNSIRIDGIT 50

51 LGGLFPVHGRGSEKPCGELKKEKGIHRLEAMLFALDRINNDPDLNPIT 100
|||||
51 LGGLFPVHGRGSEKPCGELKKEKGIHRLEAMLFALDRINNDPDLNPIT 100

101 LGARILDTCSDTHALEQSLTFVQALIEKDGTEVRCGSGGPPITTKPERV 150
|||||
101 LGARILDTCSDTHALEQSLTFVQALIEKDGTEVRCGSGGPPITTKPERV 150

151 VGVIGASGSSVSMVANILRLFKIPQISYASTAPDLSDNSRYDFFSRVVP 200
|||||
151 VGVIGASGSSVSMVANILRLFKIPQISYASTAPDLSDNSRYDFFSRVVP 200

201 SDTYQQAQAMVDIVRALKWNVYSTVASEGSYGESGVEAFIQKSREDGGVCI 250
|||||
201 SDTYQQAQAMVDIVRALKWNVYSTVASEGSYGESGVEAFIQKSREDGGVCI 250

251 AQS VKIPREP KAGEFDKIIRRLLET SNARAVIIFANEDDIRRVLEAARRA 300
|||||
251 AQS VKIPREP KAGEFDKIIRRLLET SNARAVIIFANEDDIRRVLEAARRA 300

Fig. 15

301 NQTGHEFFWGSWSKIAPVLHLEVAEGAVTILPKRMSVR..... 342
 |||||
 301 NQTGHEFFWGSWSKIAPVLHLEVAEGAVTILPKRMSVRGFDRYFSS 350
 343DRERIGQDSAY 353
 .|||
 351 RTLDNNRRNIWFAEFWEDNFHCKLSRHALKKGSHVKKCTNRERIGQDSAY 400
 354 EQEGKVQFVIDAVYAMGHALHAMHRDLCPRVGLCPRMDPVDGTQLLKYI 403
 |||||
 401 EQEGKVQFVIDAVYAMGHALHAMHRDLCPRVGLCPRMDPVDGTQLLKYI 450
 404 RNVNFSGLAGNPVTFNENGDAPEGRYDIYQYQLRNDSAEYKVIGSWTDHLH 453
 |||||
 451 RNVNFSGLAGNPVTFNENGDAPEGRYDIYQYQLRNDSAEYKVIGSWTDHLH 500
 454 LRIERMHWPGSGQQLPRSLPCQPGERKKTVKGMPCCHCEPCTGYQY 503
 |||||
 501 LRIERMHWPGSGQQLPRSLPCQPGERKKTVKGMPCCHCEPCTGYQY 550
 504 QVDRYTCKTCFYDMRPTENRTGCRPIPIIKLEWGSPWAVLPLFLAVVGIA 553
 |||||
 551 QVDRYTCKTCFYDMRPTENRTGCRPIPIIKLEWGSPWAVLPLFLAVVGIA 600

Fig. 15 (Cont.)

554 ATLFVVITFVRYNDTPIVKASGRELSYVLLAGIFLCYATTFILMIAEPDLG 603
|||||
601 ATLFVVITFVRYNDTPIVKASGRELSYVLLAGIFLCYATTFILMIAEPDLG 650

604 TCSLRRIFLGLGMSISYAALLTKTNRIYRIFEQKRSVSAPRFISPASQL 653
|||||
651 TCSLRRIFLGLGMSISYAALLTKTNRIYRIFEQKRSVSAPRFISPASQL 700

654 AITFSLISLQLLGICVWFVVDPSHSVDFQDQRTLDPRFARGVLKCDISD 703
|||||
701 AITFSLISLQLLGICVWFVVDPSHSVDFQDQRTLDPRFARGVLKCDISD 750

704 ISLICLLGYSMILLMVTCTVYAIKTRGVPETENEAKPIGFTMYTTCIVWLA 753
|||||
751 ISLICLLGYSMILLMVTCTVYAIKTRGVPETENEAKPIGFTMYTTCIVWLA 800

Fig. 15 (Cont.)

754 FIPFFGTSQSADKLYIQTTTLTVSVLSASVSLGMLYMPKVYIILFHPE 803
|||||
801 FIPFFGTSQSADKLYIQTTTLTVSVLSASVSLGMLYMPKVYIILFHPE 850
|||||
804 QNVEKRRSLKAVVTAATMSNKFTQGNFRPNGEAKSELCEALEAPALAT 853
|||||
851 QNVEKRRSLKAVVTAATMSNKFTQGNFRPNGEAKSELCEALEAPALAT 900
|||||
854 KQTYVTVYTNHAI 865
|||||
901 KQTYVTVYTNHAI 912

Fig. 15 (Cont.)

1 METKGYHSLPEGLDMERRWGQVSQAVERSLGP TERTDENNYMEIVNVSC 50
|||||
1 METKGYHSLPEGLDMERRWGQVSQAVERSLGP TERTDENNYMEIVNVSC 50
51 VSGAIPNNSTQGSSKEKQELLPCLOQDNNRPGILTS DIKTELESKELSAT 100
|||||
51 VSGAIPNNSTQGSSKEKQELLPCLOQDNNRPGILTS DIKTELESKELSAT 100
101 VAESMGLYMDSVRDADYSYEQQNQQGSMSPAKIYQNVEQLVKFYKGNCHR 150
|||||
101 VAESMGLYMDSVRDADYSYEQQNQQGSMSPAKIYQNVEQLVKFYKGNCHR 150
151 PSTLSCVNTPLRSMDSGSSVNGGVMRAIVKSPIMCHEKSPSVCPLNM 200
|||||
151 PSTLSCVNTPLRSMDSGSSVNGGVMRAIVKSPIMCHEKSPSVCPLNM 200
201 TSSVCSPAGINVSSTTASFSGFPVHSPITQGTPLT CSPNAENRGSRSHS 250
|||||
201 TSSVCSPAGINVSSTTASFSGFPVHSPITQGTPLT CSPNAENRGSRSHS 250
251 PAHASNVGSPLSPLSSMKSSISSPPSHCSVKSPVSSPNNVTLRSSVSSP 300
|||||
251 PAHASNVGSPLSPLSSMKSSISSPPSHCSVKSPVSSPNNVTLRSSVSSP 300

Fig. 16

301 ANINNSRCSVSSPSNTNRRSTLSSPAASTVGSICSPVNNAFSYTAGTSA 350
 |||||
 301 ANINNSRCSVSSPSNTNRRSTLSSPAASTVGSICSPVNNAFSYTAGTSA 350

 351 GSSTLRDVVPSPDTQEKGAQEVFPFKTEEVESAINSGVTGQLNIVQYIKP 400
 |||||
 351 GSSTLRDVVPSPDTQEKGAQEVFPFKTEEVESAINSGVTGQLNIVQYIKP 400

 401 EPDGAFASSCLGNSKINDSSFSVPIKQESTKHSCSGTSFKGNPTVNP 450
 |||||
 401 EPDGAFASSCLGNSKINDSSFSVPIKQESTKHSCSGTSFKGNPTVNP 450

 451 PFMDGSYFSEMDDKDYISLSCILGPPVPGFDGNCCEGSGFPVGIKQEPDDG 500
 |||||
 451 PFMDGSYFSEMDDKDYISLSCILGPPVPGFDGNCCEGSGFPVGIKQEPDDG 500

 501 SYYPEASIPSSAIVGVNSGGQSFHYRIGAQTISLSRSARDQSFQHLSSF 550
 |||||
 501 SYYPEASIPSSAIVGVNSGGQSFHYRIGAQTISLSRSARDQSFQHLSSF 550

 551 PPVNTLVESWKSHGDLSSRRSDGYPVLEYIPENVSSTLRSVSTGSSRPS 600
 |||||
 551 PPVNTLVESWKSHGDLSSRRSDGYPVLEYIPENVSSTLRSVSTGSSRPS 600

Fig. 16 (Cont.)

```

601 KICLVCGDEASGCHYGVTGCGCKVFFKRAVEGQHNYLCAGRNDCIIDKI 650
|||||
601 KICLVCGDEASGCHYGVTGCGCKVFFKRAVEGQHNYLCAGRNDCIIDKI 650

651 RRKNCPACRLQKCLQAGMNLGARKSKKLGKLGTHHEEQPQQQPPPPPP 700
|||||
651 RRKNCPACRLQKCLQAGMNLGARKSKKLGKLGTHHEEQPQQQPPPPPP 700

701 PQSPEEGTTYIAPAKEPSVNTALVPQLSTISRALTTPSPVMVLENIEPEIV 750
|||||
701 PQSPEEGTTYIAPAKEPSVNTALVPQLSTISRALTTPSPVMVLENIEPEIV 750

751 YAGYDSSKPDTAENLLSTLNRLAGKQMIQVVKWAKVLPGEKNLPLEDQIT 800
|||||
751 YAGYDSSKPDTAENLLSTLNRLAGKQMIQVVKWAKVLPGEKNLPLEDQIT 800

801 LIQYSWMCLSSFALSWRSYKHTNSQFLYFAPDLVFNE 837
|||||
801 LIQYSWMCLSSFALSWRSYKHTNSQFLYFAPDLVFNE 837

```

Fig. 16 (Cont.)

1 METKGYSLPEGLDMERRWGQVSQAVERSLGPRTERTDENNYMEIVNVSC 50
|||||
1 METKGYSLPEGLDMERRWGQVSQAVERSLGPRTERTDENNYMEIVNVSC 50
51 VSGAIPNNSTQGSSKEKQELLPCLOQDNNRPGILTSDIKTELESKELSAT 100
|||||
51 VSGAIPNNSTQGSSKEKQELLPCLOQDNNRPGILTSDIKTELESKELSAT 100
101 VAESMGLYMDSVRDADYSYEQNQCGSMSPAKIYQNVQVLFYKGNHR 150
|||||
101 VAESMGLYMDSVRDADYSYEQNQCGSMSPAKIYQNVQVLFYKGNHR 150
151 PSTLSCVNTPLRSEMSDSSGVNGVMRAIVKSPIMCHEKSPSVCPLNM 200
|||||
151 PSTLSCVNTPLRSEMSDSSGVNGVMRAIVKSPIMCHEKSPSVCPLNM 200
201 TSSVCSPAGINSVSSTTASEGFPVHSPITQGTPLTCSPNAENRGSRSHS 250
|||||
201 TSSVCSPAGINSVSSTTASEGFPVHSPITQGTPLTCSPNAENRGSRSHS 250
251 PAHASNVGSPLSPLSSMKSSISSPPSHCSVKSPVSSPNNVTLRSSVSSP 300
|||||
251 PAHASNVGSPLSPLSSMKSSISSPPSHCSVKSPVSSPNNVTLRSSVSSP 300

Fig. 17

301 ANINNSRCVSSPSNTNRRSTLSSPAASTVGSICSPVNNAFSYTASG TSA 350
|||||
301 ANINNSRCVSSPSNTNRRSTLSSPAASTVGSICSPVNNAFSYTASG TSA 350
351 GSSTLRDVVPSPDTQEKGAQEVFPFKTEEEVESAINSGVTGQLNIVQYIKP 400
|||||
351 GSSTLRDVVPSPDTQEKGAQEVFPFKTEEEVESAINSGVTGQLNIVQYIKP 400
401 EPDGAFFSSCLGNSKINDSSFSVPIKQESTKHSCSGTSFKGNPTVNP 450
|||||
401 EPDGAFFSSCLGNSKINDSSFSVPIKQESTKHSCSGTSFKGNPTVNP 450
451 PFMDGSYFSEMDDKDYISLGLGPPVPGFDGNCESGFPVGIKQEPDDG 500
|||||
451 PFMDGSYFSEMDDKDYISLGLGPPVPGFDGNCESGFPVGIKQEPDDG 500
501 SYYPEASIPSSAIVGVNSGGQSFHYRIGAQTISLSRSARDQSFQHLSSF 550
|||||
501 SYYPEASIPSSAIVGVNSGGQSFHYRIGAQTISLSRSARDQSFQHLSSF 550
551 PPVNTLVESWKSHGDLSSRRSDGYPVLEYIPENVSSSTLRSVSTGSSRPS 600
|||||
551 PPVNTLVESWKSHGDLSSRRSDGYPVLEYIPENVSSSTLRSVSTGSSRPS 600

Fig. 17 (Cont.)

```

601 KICLVCGDEASGCHYGVTGCGSCKVFFKRAVEGQHNYLCAGRNDCIIDKI 650
|||||
601 KICLVCGDEASGCHYGVTGCGSCKVFFKRAVEGQHNYLCAGRNDCIIDKI 650

651 RRKNCPACRLQKCLQAGMNLGARKSKKLKLGIIHEEQPQQQPPPPPP 700
|||||
651 RRKNCPACRLQKCLQAGMNLGARKSKKLKLGIIHEEQPQQQPPPPPP 700

701 PQSPEEGTYIAPAKEPSVNTALVPQLSTISRALTSPVMVLENIEPEIV 750
|||||
701 PQSPEEGTYIAPAKEPSVNTALVPQLSTISRALTSPVMVLENIEPEIV 750

751 YAGYDSSKPDTAENLLSTLNRLACKQMIQVVKWAKVLPGEKNLPLEDQIT 800
|||||
751 YAGYDSSKPDTAENLLSTLNRLACKQMIQVVKWAKVLPGEKNLPLEDQIT 800

801 LIQYSWMCLSSFALSWRSYKHTNSQELYFAPDLVENE 837
|||||
801 LIQYSWMCLSSFALSWRSYKHTNSQELYFAPDLVENE 837

```

Fig. 17 (Cont.)

```

1  MGRQLVVLGLTCCWAVASAAKLGAVYTEGGFVEGVNKKLGLLGDSDIF 50
   |||||
1  MGRQLVVLGLTCCWAVASAAKLGAVYTEGGFVEGVNKKLGLLGDSDIF 50

51  KGIPFAAPTCALENPQHPGWQGTILKAKNEKKRCLQATITQDSTYGDEDC 100
   |||||
51  KGIPFAAPTCALENPQHPGWQGTILKAKNEKKRCLQATITQDSTYGDEDC 100

101  LYLNIWVPQGRKQVSRDLPVMIWIYGGAFLMGSGHGANFLNNLYDGEI 150
   |||||
101  LYLNIWVPQGRKQVSRDLPVMIWIYGGAFLMGSGHGANFLNNLYDGEI 150

151  ATRGNVIVVTFNYRVGPLGFLSTGDANLPGNYGLRDQHMAIAWVKRNIAA 200
   |||||
151  ATRGNVIVVTFNYRVGPLGFLSTGDANLPGNYGLRDQHMAIAWVKRNIAA 200

201  FGGDPNNITLFGESAGGASVSLQTLSPYNKGLIRRAISQSGVALSPWVIQ 250
   |||||
201  FGGDPNNITLFGESAGGASVSLQTLSPYNKGLIRRAISQSGVALSPWVIQ 250

```

Fig. 18

251 KNPLFWAKKVAEKVGCPCVGDAAARMAQCLKVTDPRALTLAYKVPPLAGLEYP 300
 |||||
 251 KNPLFWAKKVAEKVGCPCVGDAAARMAQCLKVTDPRALTLAYKVPPLAGLEYP 300
 |||||
 301 MLHYVGFVPVIDGDFIPADPINLYANAADIDYIAGTNNMDGHIFASIDMP 350
 |||||
 301 MLHYVGFVPVIDGDFIPADPINLYANAADIDYIAGTNNMDGHIFASIDMP 350
 |||||
 351 AINKGNKKVTEEDFYKLVSEFTITKGLRGAKTTFDVYTESWAQDPSQENK 400
 |||||
 351 AINKGNKKVTEEDFYKLVSEFTITKGLRGAKTTFDVYTESWAQDPSQENK 400
 |||||
 401 KKTVVDFETDVLFLVPTEIALAQHRANAKSAKTYAYLFSHPSRMPVYPKW 450
 |||||
 401 KKTVVDFETDVLFLVPTEIALAQHRANAKSAKTYAYLFSHPSRMPVYPKW 450
 |||||
 451 VGADHADDIQYVFGKPFATPTGYRQDRTVSKAMIAYWNTNFAKTGDPNMG 500
 |||||
 451 VGADHADDIQYVFGKPFATPTGYRQDRTVSKAMIAYWNTNFAKTGDPNMG 500
 |||||

Fig. 18 (Cont.¹)

501 DSAVPTHWEPTYTENS GYLEITKKMGSSSMKRSLRTNFLRYWTLTYLALP 550
|||||
501 DSAVPTHWEPTYTENS GYLEITKKMGSSSMKRSLRTNFLRYWTLTYLALP 550
|||||
551 TVTDQEATPVPPPTG DSEATPVPPPTG DSETAPVPPPTG DSGAPPVPPPTG DSG 600
|||||
551 TVTDQEATPVPPPTG DSEATPVPPPTG DSETAPVPPPTG DSGAPPVPPPTG DSG 600
601 APPVPPPTG DSGAPPVPPPTG DSGAPPVPPPTG DSGAPPVPP 639
|||||
601 APPVPPPTG DSGAPPVPPPTG DSGAPPVPPPTG DSGAPPVPP 639

Fig. 18 (Cont.²)

1 MRALLARLLLCVLVSDSKGSNELHQVPSNCDCLNGGTCVSNKYFSNIHW 50
|||||
1 MRALLARLLLCVLVSDSKGSNELHQVPSNCDCLNGGTCVSNKYFSNIHW 50
51 CNCPKKFGGQHCEIDKSKTCYEGNGHFYRGKASTDTMGRPCLPWNSATVL 100
|||||
51 CNCPKKFGGQHCEIDKSKTCYEGNGHFYRGKASTDTMGRPCLPWNSATVL 100
101 QQTYHAHRSDALQLGLGKHNYCR 123
|||||
101 QQTYHAHRSDALQLGLGKHNYCR 123

Fig. 19

1 MKTYRAKFCGVCTDGRCCCTPHRTTTLPEVEFKCPDGEVMKKNNMMFIKTCAC 50
|||||
276 MKTYRAKFCGVCTDGRCCCTPHRTTTLPEVEFKCPDGEVMKKNNMMFIKTCAC 325

51 HYNCPGDNDIFESLYYRKMYGDMA 74
|||||

326 HYNCPGDNDIFESLYYRKMYGDMA 349

Fig. 20

1 MTAASMGVVRVAFVLLALCSRPAVGQNCSGPCRCPEPAPRCPAGVSLV 50
|||||
1 MTAASMGVVRVAFVLLALCSRPAVGQNCSGPCRCPEPAPRCPAGVSLV 50
51 LDGCGCCRVCAKQLGELCTERDPCDPHKGLFCDFGSPANRKIGVCTAKDG 100
|||||
51 LDGCGCCRVCAKQLGELCTERDPCDPHKGLFCDFGSPANRKIGVCTAKDG 100
101 APCIFGGTVYRSGESFQSSCKYQCTCLDGAVGCMPLCSMDVRLPSFDCPF 150
|||||
101 APCIFGGTVYRSGESFQSSCKYQCTCLDGAVGCMPLCSMDVRLPSFDCPF 150

151 P 151

151 P 151

Fig. 21

1101 NILHSRHHKPDGLYQCEASLGDGSIISRTAKVAVAGP 139
 1101 NILHSRHHKPDGLYQCEASLGDGSIISRTAKVAVAGP 139

Fig. 22

```

1 MAPFEPLASGILLLLWLIAPSRACCTVPPHPQTAFNCNSDLVIRAKFVGTP 50
  |||||
1 MAPFEPLASGILLLLWLIAPSRACCTVPPHPQTAFNCNSDLVIRAKFVGTP 50
  |||||
51 EVNQTTLYQRYEIKMTKMYKGFQALXDAADIRFVYTPAMESVCCGYFHRSH 100
  |||||
51 EVNQTTLYQRYEIKMTKMYKGFQALGDAADIRFVYTPAMESVCCGYFHRSH 100
  |||||

101 NRS 103
  |||
101 NRS 103

```

Fig. 23

13 QMYKGFQALGDAADIRFVYTPAMESVCGYFHRSHNRSEEFLLIAGE 57
 .|||||
 67 KMYKGFQALGDAADIRFVYTPAMESVCGYFHRSHNRSEEFLLIAGK 111

Fig. 24


```

1  MAPFEPLASGILLLLWLIAPSRACTCVPPHPQTAFCNSDLVIRAKFVGTP 50
      .  |||||  |||||  |||||  |||||  |||||  |||||  |||||  |||||  |||||
1  MAPFEPLASGILLLLWLIAPSRACTCVPPHPQTAFCNSDLVIRAKFVGTP 50

51  EVNQTTLYQRYEIKMTKMYKGFQALGDAADIRFVYTPAMESVCGYFHR 98
      .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .  .
51  EVNQTTLYQRYEIKMTKMYKGFQALGDAADIRFVYTPAMESVCGYFHR 98

```

1 MTKMYKGFQALGDAADIRFVYTPAMESVCGYFHRSHNRSEEFLLIAGKLQD 50
|||||
65 MTKMYKGFQALGDAADIRFVYTPAMESVCGYFHRSHNRSEEFLLIAGKLQD 114
51 GLLHITTCSFVAPWNSLSLAQRRGFTKTYTVGCCEECTVFPCLSI[.]PCCKLQS 100
|||||
115 GLLHITTCSFVAPWNSLSLAQRRGFTKTYTVGCCEECTVFPCLSI[.]PCCKLQS 164
101 GTHCLWTDQLLQSEKGFQSRHLACLPREPGLCTWQSLRSQIA 143
|||||
165 GTHCLWTDQLLQSEKGFQSRHLACLPREPGLCTWQSLRSQIA 207

Fig. 26